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TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371

09/509482

INTERNATIONAL APPLICATION NO. PCT/AU98/00817

INTERNATIONAL FILING DATE 29 September 1998

PRIORITY DATE CLAIMED

29 September 1997

TITLE OF INVENTION

Isoforms of the Human Vitamin D Receptor

APPLICANT(S) FOR DO/EO/US

Linda Anne CROFTS, Manuella S. HANCOCK, Nigel A. MORRISON, John A. EISMAN

Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:

- . [X] This is a FIRST submission of items concerning a filing under 35 U.S.C. 371
- 2. This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371.
- 3. [] This express request to begin national examination procedures (35 U.S.C. 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39(1).
- 4. [X] A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date.
- 5. [X] A copy of the International Application as filed (35 U.S.C. 371(c)(2))
 - a. [] is transmitted herewith (required only if not transmitted by the International Bureau).
 - b. [X] has been transmitted by the International Bureau.
 - c. [] is not required, as the application was filed in the United States Receiving Office (RO/US)
- 6. [] A translation of the International Application into English (35 U.S.C. 371(c)(2)).
- 7. [X] Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3))
 - a. [] are transmitted herewith (required only if not transmitted by the International Bureau).
 - b. [] have been transmitted by the International Bureau.
 - c. [] have not been made; however, the time limit for making such amendments has NOT expired.
 - d. [X] have not been made and will not be made.
- 3. A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).
- An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).
- A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(e)(5)).

ITEMS 11. TO 16. below concern other document(s) or information included:

- 11. [X] An Information Disclosure Statement under 37 CFR 1.97 and 1.98.
- 12. [] An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
- 13. [X] A FIRST preliminary amendment.
 - A SECOND or SUBSEQUENT preliminary amendment.
- 14. [] A substitute specification.
- 15. [] A change of power of attorney and/or address letter.
- 16. [X] Other items or information:
 - Copy of published application 99/16872
 - Copy of International Preliminary Examination Report
 - Paper copy and computer copy of sequence listing

122 Rec'd PCT/PTO 2 9 MAR 2000

U.S. APPLICATION NO. (If known, s	ATTORNEY DOCKET NO. 1871–130										
17. [X] The following Basic Nations Search Report has beer International prelimina No international prelim but international search Neither international search International prelimina and all claims satisfied	CALCULATIONS	PTO USE ONLY									
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Claims	Claims Number Filed Number Extra Rate										
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Independent Claims	6 - 3 =	3	X \$78.00	\$ 234.00							
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b. [] Please charge my Deposit Account No. 02-2135 in the amount of \$ to cover the above fees. A duplicate copy of this sheet is enclosed.											
c. [X] The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 02-2135. A duplicate copy of this sheet is enclosed.											
NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.											
SEND ALL CORRESPONI Barbara G. Ernst Rothwell, Figg, Ernst & Ku 555 13th St., N.W. Washington, D.C. 20004 Phone: 202/783-6040	Signature Barbara G. Ernst Name 30,377	- G Ernot									
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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of)
Linda Anne CROFTS et al.)) Filing Under 35 USC 371) International Application
Serial No.) No. PCT/AU98/00817
Filed:) Filed 29 September 1998)
For: ISOFORMS OF THE HUMAN VITAMIN D RECEPTOR)))

PRELIMINARY AMENDMENT

Assistant Commissioner for Patents Washington, D.C. 20231

Dear Sir:

Prior to calculating the filing fee for the above-referenced patent application, please enter the following amendments:

In the Claims:

In claim 9, line 2, please delete "any one of the preceding claims" and insert therefor --claim 1--.

In claim 10, line 2, please delete "any one of claims 1-8" and insert therefor --claim 1 or 5--.

In claim 10, line 2, at the end of the line, please insert - or 25--.

In claim 13, lines 2-3, please delete "any one of claims 10-12" and insert therefor --claim 10--.

In claim 15, line 3, please delete "any one of claims 1-4" and insert therefor --claim 1--.

In claim 17, line 2, please delete "any one of claims 1-8" and insert therefor --claim 1 or 5--.

In claim 18, line 4, please delete "any one of claims 1-4" and insert therefor --claim 1--.

In claim 19, line 4, please delete "any one of claims 1-8" and insert therefor --claim 1 or 5--.

In claim 20, line 4, please delete "any one of claims 1-8" and insert therefor --claim 1 or 5--.

Please add the following new claim:

--25. A plasmid or expression vector including a polynucleotide molecule according to claim 5.--

REMARKS

The amendments set forth above are made to correct improper multiple dependent claims. No new matter is introduced into the application through these amendments.

Respectfully submitted,

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Field of the Invention:-

The present invention relates to isolated polynucleotide molecules which encode novel isoforms of the human Vitamin D receptor (hVDR) or variant transcripts for hVDR. The polynucleotide molecules may be utilised in, for example, methods of screening compounds for VDR agonists and/or antagonists.

10 Background of the Invention:-

The active hormonal form of vitamin D, 1,25-dihydroxyvitamin D₃ (1.25(OH)₂D₃), has a central role in calcium and phosphate homeostasis, and the maintenance of bone. Apart from these calcitropic effects. 1.25-(OH)₂D₃ has been shown to play a role in controlling cell growth and differentiation in many target tissues. The effects of 1,25-(OH)₂D₃ are mediated by a specific receptor protein, the vitamin D receptor (VDR), a member of the nuclear receptor superfamily of transcriptional regulators which also includes steroid, thyroid and retinoid receptors as well as a growing number of orphan receptors. Upon binding hormone the VDR regulates gene expression by direct interaction with specific sequence elements in the promotor regions of hormone responsive target genes. This transactivation or repression involves multiple interactions with other protein cofactors, heterodimerisation partners and the transcription machinery.

Although a cDNA encoding the human VDR was cloned in 1988 (1), little has been documented characterising the gene structure and pattern of transcription since that time. The regulation of VDR abundance is one potentially important mechanism for modulating 1.25-(OH)₂D₃ responsiveness in target cells. It is also possible that VDR has a role in non-transcriptional pathways, perhaps via localization to a non-nuclear compartment and/or interaction with components of other signalling pathways. However, the question of how VDRs are targetted to different cell types and how they are regulated remains unresolved. There have been many reports in the literature describing translational or transcriptional control of VDR levels, both homologously and heterologously, mostly in non-human systems.

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A recent study (2) showed that in the kidney, alternative splicing of human VDR transcripts transcribed from a GC rich promotor generates several transcripts which vary only in their 5' UTRs. The present inventors have now identified further upstream exons of the VDR gene which generate 5' variant transcripts, suggesting that the expression of the VDR gene is regulated by more than one promoter. A subset of these transcripts is expressed in a restricted tissue-specific pattern and further variant transcripts have the potential to encode an N-terminally variant protein. These results may have implications for understanding the actions of 1,25-(OH)₂D₃ in different tissues and cell types, and the possibility that N-terminally variant VDR proteins may be produced has implications for altered activities such as transactivation function or subcellular localisation of the receptor protein. Furthermore, these variants, by their level, tissue specificity, subcellular localisation and functional activity, may yield targets for pharmaceutical intervention. The variants may also be useful in screening potential analogs and/or antagonists of vitamin D compounds.

Disclosure of the Invention:-

In a first aspect, the invention provides an isolated polynucleotide molecule encoding a human Vitamin D receptor (hVDR) isoform, said polynucleotide molecule comprising a nucleotide sequence which includes sequence that substantially corresponds or is functionally equivalent to that of exon 1d of the human VDR gene.

Exon 1d (referred to as exon 1b in the Australian Provisional Patent Specification No. PO9500) is a 96 bp exon located 296 bp downstream from exon 1a (2). The sequence of exon 1d is:

5'GTTTCCTTCTGTCGGGGCGCCCTTGGCATGGAGTGGAGGAATAAGAA AAGGAGCGATTGGCTGTCGATGGTGCTCAGAACTGCTGGAGTGGAGG3' (SEQ ID NO: 1).

The nucleotide sequence of the polynucleotide molecule of the first aspect of the invention, preferably does not include sequence corresponding to that of exon 1a, exon 1f and/or exon 1e. However, the nucleotide sequence of the polynucleotide molecule of the first aspect of the invention, may or

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may not include sequence that substantially corresponds or is functionally equivalent to that of exon 1b and/or exon 1c.

Preferably, the polynucleotide molecule of the first aspect comprises a nucleotide sequence which includes;

- (i) sequence that substantially corresponds or is functionally equivalent to that of exons 1d, 1c and 2-9 and encodes a VDR isoform of approximately 477 amino acids,
- (ii) sequence that substantially corresponds or is functionally equivalent to that of exons 1d and 2-9 and encodes a VDR isoform of approximately 450 amino acids, or
- (iii) sequence that substantially corresponds or is functionally equivalent to that of exons 1d and 2-9 and further includes a 152 bp intronic sequence, and encodes a truncated VDR isoform of approximately 72 amino acids.

Most preferably, the polynucleotide molecule of the first aspect of the invention comprises a nucleotide sequence substantially corresponding to that shown as SEQ ID NO: 2, SEQ ID NO: 3 or SEQ ID NO: 4.

In a second aspect, the invention provides an isolated polynucleotide molecule encoding a human Vitamin D receptor (hVDR), said polynucleotide molecule comprising a nucleotide sequence which includes sequence that substantially corresponds to that of exon 1f and/or 1e of the human VDR gene.

Exon 1f is a 207bp exon located more than 9kb upstream from exon 1a (2) bp upstream from exon 1c(8). The sequence of exon 1f is:

Exon 1e is a 157 bp exon located 1826bp upstream from exon 1a (2). The sequence of exon 1e is:

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5'AGGCAGCATGAAACAGTGGGATGTGCAGAG
AGAAGATCTGGGTCCAGTAGCTCTGACACTCCTCAGCTGT
AGAAACCTTGACAACTCTGCACATCAGTTGTACAATGGAA
CGGTATTTTTTACTCTTCATGTCTGAAAAGGCTATGATAA
AGATCAA3' (SEQ ID NO: 6)

The nucleotide sequence of the polynucleotide molecule of the second aspect of the invention, preferably does not include sequence corresponding to that of exon 1a, 1d or 1b. However, the nucleotide sequence of the polynucleotide molecule of the second aspect of the invention, may or may not include sequence that substantially corresponds or is functionally equivalent to that of exon 1c.

Preferably, the nucleotide molecule of the second aspect comprises a nucleotide sequence which includes sequence that substantially corresponds or is functionally equivalent to that of exons 1f and 2-9.

Most preferably, the polynucleotide molecule of the first aspect of the invention comprises a nucleotide sequence substantially corresponding to that shown as SEQ ID NO: 7.

The polynucleotide molecule of the first or second aspects may be incorporated into plasmids or expression vectors (including viral vectors), which may then be introduced into suitable host cells (e.g. bacterial, yeast, insect and mammalian host cells). Such host cells may be used to express the VDR or functionally equivalent fragment thereof encoded by the isolated polynucleotide molecule.

Accordingly, in a third aspect, the present invention provides a host cell transformed with the polynucleotide molecule of the first or second aspect.

In a fourth aspect, the present invention provides a method of producing a VDR or a functionally equivalent fragment thereof, comprising culturing the host cell of the first or second aspect under conditions enabling the expression of the polynucleotide molecule and, optionally, recovering the VDR or functionally equivalent fragment thereof.

Preferably, the host cell is of mammalian origin. Preferred examples include NIH 3T3 and COS 7 cells.

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In a preferred embodiment, the VDR or functionally equivalent fragment thereof is localised to a cell membrane or other subcellular compartment as distinct from a nuclear localisation.

The polynucleotide molecules of the first aspect of the invention encode novel VDR isoforms which may be of interest both clinically and commercially. By using the polynucleotide molecule of the present invention it is possible to obtain VDR isoform proteins or functionally equivalent fragments thereof in a substantially pure form.

Accordingly, in a fifth aspect, the present invention provides a human VDR isoform or functionally equivalent fragment thereof encoded by a polynucleotide molecule of the first aspect, said VDR isoform or functionally equivalent fragment thereof being in a substantially pure form.

In a sixth aspect, the present invention provides an antibody or antibody fragment capable of specifically binding to the VDR isoform of the fourth aspect.

The antibody may be monoclonal or polyclonal, however, it is presently preferred that the antibody is a monoclonal antibody. Suitable antibody fragments include Fab, F(ab')₂ and scFv.

In an eighth aspect, the present invention provides a non-human animal transformed with a polynucleotide molecule according to the first or second aspect of the invention.

In a seventh aspect, the invention provides a method for detecting agonist and/or antagonist compounds of a VDR isoform of the fourth aspect, comprising contacting said VDR isoform, functionally equivalent fragment thereof or a cell transformed with and expressing the polynucleotide molecule of the first aspect, with a test compound under conditions enabling the activation of the VDR isoform or functionally equivalent fragment thereof, and detecting an increase or decrease in the activity of the VDR isoform or functionally equivalent fragment thereof.

An increase or decrease in activity of the receptor or functionally equivalent fragment thereof may be detected by measuring changes in interactions with known cofactors (e.g. SRC-1, GRIP-1 and TFIIB) or unknown cofactors (e.g. through use of the yeast dual hybrid system).

In a ninth aspect, the present invention provides an oligonucleotide or polynucleotide probe comprising a nucleotide sequence of 10 or more nucleotides, the probe comprising a nucleotide sequence such that the probe

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specifically hybridises to the polynucleotide molecule of the first or second aspect under high stringency conditions (Sambrook et al., Molecular Cloning: a laboratory manual, Second Edition, Cold Spring Harbor Laboratory Press).

Preferably, the probe is labelled.

In a tenth aspect, the present invention provides an antisense polynucleotide molecule comprising a nucleotide sequence capable of specifically hybridising to an mRNA molecule which encodes a VDR encoded by the polynucleotide molecule of the first or second aspect, so as to prevent translation of the mRNA molecule.

Such antisense polynucleotide molecules may include a ribozyme region to catalytically inactivate mRNA to which it is hybridised.

The polynucleotide molecule of the first or second aspect of the invention may be a dominant negative mutant which encodes a gene product causing an altered phenotype by, for example, reducing or eliminating the activity of endogenous VDR.

In an eleventh aspect, the invention provides an isolated polynucleotide molecule comprising a nucleotide sequence substantially corresponding or, at least, showing >75% (preferably >85% or, even more preferably, >95%) sequence identity to:

- (ii) 5'AGGCAGCATGAAACAGTGGGATGTGCAGAGAGAAGATCTGGGTC CAGTAGCTCTGACACTCCTCAGCTGTAGAAACCTTGACAACTCTGCACAT CAGTTGTACAATGGAACGGTATTTTTTACTCTTCATGTCTGAAAAAGGCTA TGATAAAGATCAA3' (exon 1e) (SEQ ID NO: 6), or
- (iii) 5'GTTTCCTTCTGTCGGGGCGCCCTTGGCATGGAGTGGAGGAATA AGAAAAGGAGCGATTGGCTGTCGATGGTGCTCAGAACTGCTGGAGTGGA GG3' (exon 1d) (SEQ ID NO: 1).

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The polynucleotide molecules of the eleventh aspect may be useful as probes for the detection of VDR variant transcripts and as such may be useful in assessing cell or tissue-specific expression of variant transcripts.

The terms "substantially corresponds" and "substantially corresponding" as used herein in relation to nucleotide sequences is intended to encompass minor variations in the nucleotide sequence which due to degeneracy in the DNA code do not result in a substantial change in the encoded protein. Further, this term is intended to encompass other minor variations in the sequence which may be required to enhance expression in a particular system but in which the variations do not result in a decrease in biological activity of the encoded protein.

The term "functionally equivalent" as used herein in relation to nucleotide sequences encoding a VDR isoform is intended to encompass nucleotide sequence variants of up to 5% sequence divergence (i.e. retaining 95% or more sequence identity) which encode VDR isoforms of substantially equivalent biological activity(ies) as said VDR isoform.

The term "functionally equivalent fragment" as used herein in respect of a VDR isoform is intended to encompass functional peptide and polypeptide fragments of said VDR isoform which include the domain or domains which bestow the biological activity characteristic of said VDR isoform.

The terms "comprise". "comprises" and "comprising" as used throughout the specification are intended to refer to the inclusion of a stated step, component or feature or group of steps, components or features with or without the inclusion of a further step, component or feature or group of steps, components or features.

The invention will hereinafter be further described by way of the following non-limiting example and accompanying figures.

Brief description of the figures:-

FIG.1. (A) Human VDR gene locus. Four overlapping cosmid clones were isolated from a human lymphocyte genomic library (Stratagene) and directly sequenced. Clone J5 extends from the 5' flanking region to intron 2; AE, from intron 1b to intron 5; D2, from intron 3 to the 3' UTR: WE, from intron 6 through the 3' flanking region. Sequence upstream of exon 1f was obtained by

anchored PCR from genomic DNA. (B) Structure of hVDR transcripts. Transcripts 1–5 originate from exon 1a. Transcript 1 corresponds to the published cDNA (1). Transcripts 6–10 originate from exon 1d and transcripts 11–14 originate from exon 1f. Boxed numbers indicate the major transcript (based on the relative intensities of the multiple PCR products) within each exon-specific group of transcripts generated with a single primer set. While all transcripts have a translation initiation codon in exon 2, exon 1d transcripts have the potential to initiate translation upstream in exon 1d, with transcripts 6 and 9 encoding VDR proteins with extended N termini. (C) N-terminal variant proteins encoded by novel hVDR transcripts. Transcript 1 corresponds to the published cDNA sequence (1) and encodes the 427-aa hVDR protein. Transcripts 6 and 9 code for a protein with an extra 50 aa or 23 aa, respectively, at the N-terminal. The 23 aa of the hVDR A/B domain are shown in bold.

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FIG. 2. RT-PCR analysis of expression of variant hVDR transcripts. (A) Exon 1a transcripts (220 bp. 301 bp, 342 bp, 372 bp, and 423 bp). (B) Exon 1d transcripts (224 bp, 305 bp, 346 bp, 376 bp. and 427 bp). (C) Exon 1f transcripts (228 bp. 309 bp. 387 bp, and 468 bp). RT-PCR was carried out with exon 1a-, 1d-. or 1f-specific forward primers and a common reverse primer in exon 3. The sizes of the PCR products and the pattern of bands are similar in A and B by virtue of the identical splicing pattern of exon 1a and 1d transcripts and the fact that primers were designed to generate PCR products of comparable sizes. All tissues and cell lines are human in origin.

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FIG. 3. Functional analysis of sequence-flanking exons 1a and 1d (A) and exon 1f (B) in NIH 3T3 (solid bars) and COS 7 cells (open bars). The parent vector pGL3basic was used as a promoterless control, and a promoter-chloramphenical acetyltransferase (CAT) gene reporter construct was cotransfected as an internal control for transfection efficiency in each case. The activity of each construct was corrected for transfection efficiency and for the activity of the pGL3basic empty vector control and expressed as a percentage of the activity of the construct 1a(-488,+75) SEM of at least three separate transfections. Exon 1a and 1d flanking constructs are defined in relation to the transcription start site of exon

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1a, designated 11, which lies 54 nt upstream of the published cDNA (1). Exon 1f flanking constructs are defined relative to the exon 1f transcription start site, designated 11. Transcription start sites were determined by the 5' termini of the longest RACE clones. The open box corresponds to the GC-rich region.

FIG 4. Provides the nucleotide sequence of novel exons detected by 5' RACE: (A) exon 1b (SEQ ID NO: 8), (B) exon 1f (SEQ ID NO: 5) [P1f is indicated by an arrow above the sequence], (C) exon 1e (SEQ ID NO: 6), (D) exon 1d (SEQ ID NO: 1) [in-frame ATG codons are highlighted and P1d is indicated by an arrow above the sequence]. Intronic sequences are shown in lower case. Canonical splice site consensus sequences are indicated in bold. The transcription start sites for exons 1f and 1d were determined by the 5' termini of RACE clones. No intron sequence is shown 3' to exon 1f as cosmid clone 15 terminated in the intron between exons 1f and 1e.

FIG 5. Provides the nucleotide sequence corresponding to transcript 6 (see figure 1) (SEQ ID NO: 2), together with the predicted amino acid sequence (SEQ ID NO: 9) of the encoded protein. Nucleotides 1-96 correspond to exon 1d; nucleotides 97-1463 correspond to exons 1c to the stop codon in exon 9 (or nucleotides -83-1283 of the hVDR cDNA (1)).

FIG 6. Provides the nucleotide sequence corresponding to transcript 9 (see figure 1) (SEQ ID NO: 3), together with the predicted amino acid sequence (SEQ ID NO: 10) of the encoded protein. Nucleotides 1-96 correspond to exon 1d; nucleotides 97 - 1382 correspond to exon 2 to the stop codon in exon 9 (or nucleotides -2 - 1283 of the hVDR cDNA (1)).

FIG 7. Provides the nucleotide sequence corresponding to transcript 10 (see figure 1) (SEQ ID NO: 4), together with the predicted amino acid sequence (SEQ ID NO: 11) of the encoded protein. Nucleotides 1-96 correspond to exon 1d; nucleotides 97-244 correspond to exon 2; nucleotides 245-396 correspond to intronic sequence immediately 3' to exon 2; nucleotides 397-1534 correspond to exons 3 to the stop codon in exon 9 (or nucleotides 146-1283 of the hVDR cDNA (1)).

FIG 8. Provides the nucleotide sequence corresponding to transcript 11 (see figure 1) (SEQ ID NO: 7), together with the predicted amino acid sequence (SEQ ID NO: 12) of the encoded protein. Nucleotides 1-207 correspond to exon 1f; nucleotides 208-1574 correspond to exon 1c to the stop codon in exon 9 (or nucleotides -83-1283 of the hVDR cDNA (1)).

Example:-

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EXPERIMENTAL PROCEDURES

Isolation and Characterisation of Genomic Clones

A human lymphocyte cosmic library (Stratagene, La Jolla, Ca) was screened using a 2.1kb fragment of the hVDR cDNA encompassing the entire coding region but lacking the 3'UTR, a 241 bp PCR product spanning exons 1 to 3 of the human VDR cDNA, and a 303 bp PCR product spanning exons 3 and 4 of the hVDR cDNA, following standard colony hybridisation techniques. DNA probes were labelled by nick translation (Life Technologies, Gaithersburg, MD) with $[\alpha^{32} P]$ dCTP. Positively hybridising colonies were picked and secondary and tertiary screens carried out until complete purification. Cosmid DNA from positive clones was purified (Qiagen), digested with different restriction enzymes and characterised by Southern blot analysis using specific $[\gamma^{32}]$ PATP labelled oligonucleotides as probes. Cosmid clones were directly sequenced using dye-termination chemistry and automated fluorescent sequencing on an ABI Prism. 377 DNA Sequencer (Perkin-Elmer, Foster City, Ca). Sequence upstream of the most 5' cosmid was obtained by anchored PCR from genomic DNA using commercially available anchor ligated DNA (Clontech, Palo Alto, Ca).

Rapid Amplification of cDNA 5-prime Ends (5'-RACE)

Alternative 5' variants of the human VDR gene were identified by 5'RACE using commercially prepared anchor-ligated cDNA (Clontech) following the instructions of the manufacturer. Two rounds of PCR using nested reverse primers in exons 3 and 2 (P 1: 5'ccgcttcatgcttcgcctgaagaagcc-3', P2: 5'-tgcagaattcacaggtcatagcattgaag-3') were carried out on a Corbett FTS-4000 Capillary Thermal Sequencer (Corbett Research, NSW, Australia). After 26 cycles of PCR, 2% of the primary reaction was reamplified for 31 cycles.

The PCR products were cloned into PUC18 and sequenced by the dideoxy chain termination method.

Cell-Culture

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The embryonal kidney cell line, HEK-293, an embryonic intestine cell line, Intestine-407 and WS 1, a foetal skin fibroblast cell line were all cultured in Eagle's MEM with Earle's BSS and supplemented with either 10% heat-inactivated FBS, 15% FBS or 10% FBS with non-essential amino acids, respectively. The osteosarcoma cell lines MG-63 and Saos-2 were cultured in Eagle's MEM with nonessential amino acids and 10% heat-inactivated FBS and McCoy's 5a medium with 15% FBS, respectively. The breast carcinoma cell line T47D and the colon carcinoma cell lines LIM 1863 and COLO 206F were cultured in RPMI medium supplemented with 0.2 IU bovine insulin/ml and 10% FBS, 5% FBS or 10% FBS, respectively. LIM 1863 were a gift from R.H. Whitehead (3). HK-2 kidney proximal tubule cells were grown in keratinocyte-serum free medium supplemented with 5ng/ml recombinant EGF, 40ug/ml bovine pituitary extract. BC1 foetal osteoblast-like cells were kindly donated by R. Mason (4) and were grown in Eagle's MEM with 5% FBS and 5mg/L vitamin C. Unless otherwise stated all cell lines were obtained from the American Type Culture Collection (Manassas, VA).

Reverse Transcriptase-PCR (RT-PCR).

Total RNA extracted from approximately 1.5 x 10³ cells. from leukocytes prepared from 40 ml blood, or from human tissue using acidphenol extraction was purified by using a guanidium isothiocyanate-cesium 25 chloride step gradient. First-strand cDNA was synthesized from 5 µg of total RNA primed with random hexamers (Promega) using Superscript II reverse transcriptase (Life Technologies). One-tenth of the cDNA (2µl) was used for subsequent PCR, with 36 cycles of amplification, using exon-specific forward primers (exon 1a: corresponding to nucleotides 1-21 of hVDR cDNA (1); 30 exon 1d: 5'-GGCTGTCGATGGTGCTCAGAAC-3'; exon 1f: 5'-AAGTTCCTCCGAGGAGCCTGCC-3'); and a common reverse primer in exon 3 [corresponding to nucleotides 301– 280 of hVDR cDNA (1)]. All RT-PCRs were repeated multiple times by using RNA/cDNA prepared at different times from multiple sources. Each PCR 35 included an appropriate cDNA-negative control, and additional controls

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included RT-negative controls prepared alongside cDNA and RNA/cDNA prepared from VDR-negative cell lines. PCR products were separated on 2% agarose and visualized with ethidium bromide staining.

5 Functional Analysis of hVDR Gene Promoters.

Sequences flanking exons 1a, 1d, and 1f (see Fig. 1A) were PCRamplified by using Pfu polymerase (Stratagene) and cloned into the pGL3basic vector (Promega) upstream of the luciferase gene reporter. Promoter-reporter constructs were transfected into NIH 3T3 and COS 7 cells by using the standard calcium phosphate-precipitation method. Cells were seeded at 2.3±2.5 x 10⁶ per 150-cm² flask the day before transfection. Several hours before the precipitates were added the medium was changed to DMEM with 2% charcoal-stripped FBS. Cells were exposed to precipitate for 16 h before subculturing and were harvested 24 h later. The parent vector pGL3basic was used as a promoterless control in these experiments and a simian virus 40 promoter-chloramphenicol acetyltransferase (CAT) gene reporter construct was cotransfected as an internal control for transfection efficiency in each case. The activity of each construct was corrected for transfection efficiency and for the activity of the pGL3 basic empty vector control and expressed as a percentage of the activity of the construct 1a(-488, +75). Luciferase and CAT assays were carried out in triplicate, and each construct was tested in transfection at least three times.

RESULTS

Identification of Alternative 5' Variants of the hVDR Gene.

Upstream exons were identified in human kidney VDR transcripts by 5' RACE (exons 1f, 1e, 1d. and 1b) and localized by sequencing of cosmid clones (Fig. 1A). To verify these results and to characterize the structure of the 5' end of the VDR gene, exon-specific forward primers were used with a common reverse primer in exon 3 to amplify specific VDR transcripts from human tissue and cell line RNA (Fig. 1B). The identity of these PCR products was verified by Southern blot and by cloning and sequencing. Five different VDR transcripts originating from exon 1a were identified. The major transcript (transcript 1 in Fig. 1B) corresponds to the published cDNA sequence (1). Three less-abundant forms (2, 3, and 4 in Fig. 1B) arise from

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alternative splicing of exon 1c and a novel 122-bp exon 1b into or out of the final transcript. These three variant transcripts were described recently by Pike and colleagues (2). A fifth minor variant was identified (5 in Fig. 1B) that lacks exons 1b and 1c, but includes an extra 152 bp of intronic sequence immediately 3' to exon 2, potentially encoding a truncated protein as a result of an in-frame termination codon in intron 2.

Four more transcripts were characterized that originate from exon 1f, a novel 207-bp exon more than 9 kb upstream from exon 1a. The major 1f-containing transcript (11 in Fig.1B) consists of exon 1f spliced immediately adjacent to exon 1c. Three less-abundant variants (12, 13, and 14 in Fig. 1B) arise from alternative splicing of exon 1c and a novel 159-bp exon 1e into or out of the final transcript. All these hVDR variants differ only in their 5' UTRs and encode identical proteins from translation initiation in exon 2.

Of considerable interest, another five hVDR transcripts were identified that originate from exon 1d, a novel 96-bp exon located 296 bp downstream from exon 1a. The major exon 1d-containing transcript (6 in Fig. 1B) utilizes exon 1d in place of exon 1a of the hVDR cDNA. Three minor variants (7, 8, and 9 in Fig. 1B) arise from alternative splicing of exons 1b and 1c into or out of the transcript, analogous to the exon 1a-containing variants 2, 3, and 4. A fifth minor variant transcript (10 in Fig. 1B) lacks exons 1b and 1c, but includes 152 bp of intron 2 analogous to the exon 1a-containing transcript 5, and also potentially encodes a truncated protein. Two of these exon 1d-containing hVDR transcripts encode an N-terminal variant form of the hVDR protein. Utilization of an ATG codon in exon 1d, which is in a favorable context and in-frame with the major translation start site in exon 2, would generate a protein with an additional 50 aa N-terminal to the ATG codon in exon 2 in the case of variant 6 or 23 aa in the case of variant 9 (Fig. 1C).

The relative level of expression of the different transcripts is difficult to address with PCR since relatively minor transcripts may be amplified. However, Southern blots of PCR products from the linear range of PCR amplification indicated that equivalent amounts of PCR product were accumulated after 26 cycles for exon 1a transcripts compared with 30 cycles for exon 1d transcripts. suggesting that 1d abundance is about 5% of that of 1a transcripts. This is consistent with the frequency of clones selected and sequenced from RACE analysis of two separate samples of kidney RNA: 1a (21/27;78%), 1d (2/27; 7%), and 1f (4/27: 15%). RT-PCR with exon 1a-, 1d-, or

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1f-specific forward primers and reverse primers in exons 7 or 9, followed by cloning and sequencing, suggests that these 5' variant transcripts are not associated with differences at the 3' end of the transcript.

5 Exon-Intron Organization of the hVDR Gene.

Overlapping cosmid clones were isolated from a human lymphocyte genomic library and characterized by hybridization to exon-specific oligonucleotide probes (Fig. 1A). The exon-intron boundaries of the hVDR gene were determined by comparison of the genomic sequence from cosmid clones with the cDNA sequence. Upstream exons were localized in the VDR gene by sequencing cosmid clones, which extend approximately 7 kb into the intron between exons 1e and 1f, enabling verification of both their sequence and the presence of consensus splice donor/acceptor sites. Sequence upstream of exon 1f was obtained by anchored PCR from genomic DNA by using commercially available anchor-ligated DNA (CLONTECH). In total, the hVDR gene spans more than 60 kb and consists of at least 14 exons (Fig. 1A).

Tissue-Specific Expression of hVDR Transcripts.

The pattern of expression of variant hVDR transcripts was examined by RT-PCR in a variety of cell lines and tissues with exon 1a-, 1d-, or 1f-specific forward primers and a common reverse primer in exon 3. Exon 1a and 1d transcripts (Fig. 1B, variants 1–10) were coordinately expressed in all RNA samples analyzed (Fig. 2 A and B). Exon 1f transcripts (Fig. 1B, variants 11–14), however, were detected only in RNA from human kidney tissue (two separate samples), human parathyroid adenoma tissue, and an intestinal carcinoma cell line, LIM 1863 (Fig. 2C). Interestingly, these represent major target tissues for the calcitropic effects of vitamin D.

Functional Analysis of hVDR Gene Promoters.

Promoter activities of the 5' flanking regions of exons 1a, 1d, and 1f were examined in NIH 3T3 and COS 7 cells (Fig. 3). Sequences flanking exon 1a exhibited high promoter activity in both cell lines (Fig. 3A). Maximum luciferase expression of 36- and 54-fold over the empty vector was attained for construct 1a(-488,+75) in NIH 3T3 and COS 7 cells, respectively. This activity could be attributed largely to a GC-rich region containing multiple consensus Sp1-binding motifs lying within 100 bp immediately adjacent to

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the transcription start site. This region alone, upstream of a luciferase reporter [construct 1a(-94.+75)], accounted for 43% of the maximum activity observed in NIH 3T3 cells and 86% of the maximum observed in COS 7 cells. The removal of this GC-rich region [construct 1a(-29.+75)] reduced luciferase activity to only 13% of the maximum in NIH 3T3 and 19% in COS 7 cells. Despite the fact that VDR transcripts that originated from exon 1d were identified, distinct promoter activity was not associated with sequences within 300 bp of exon 1d [constructs 1d(+87,+424) and 1d(+244,+424)]; rather, the sequence immediately adjacent to exon 1d may contain a suppressor element (Fig. 3A). Construct 1a-1d(-846,+470), spanning the 5' flanking regions of both exons 1a and 1d. resulted in only 42% and 60% of the activity of 1a(-898.+75) in NIH 3T3 and COS 7 cells, whereas the 3'deletion of 227 bp restored luciferase activity to 65% and 97% of the activity of 1a(-898.+75), respectively. Similarly. the 5' truncated construct 1a-1d (-94,+470), spanning the 5' flanking regions of both 1a and 1d. resulted in only 35% and 40% of the activity of 1a(-94, +75), while a further 3' deletion of 227bp restored luciferase activity to 69% and 91% of the activity of 1a(-94, +75)in NIH 3T3 and COS 7 cells. It is possible that transcription from exons 1a and 1d is driven by overlapping promoter regions rather than from two distinct promoters, as has been described for the mouse androgen receptor gene.

Sequence upstream of exon 1f showed significant promoter activity in NIH 3T3 cells of 22% of that of the most active construct, 1a(-488,+75), or 9-fold over pGL3basic [construct 1f(-1168,+58)] (Fig. 3B). A shorter construct [1f(-172,+58)] had similar activity, with evidence of a suppressor element (between nucleotides -278 and +172) able to repress luciferase activity by 70%. Interestingly, the same constructs were not active in COS 7 cells. This cell line-specific activity of exon 1f flanking sequences may reflect a requirement for tissue- or cell-specific protein factors.

Identification of VDR isoforms in whole cell lysates

The existence of a VDR isoform including exons 1d and 1c has been confirmed in cell lysates from multiple human, monkey, rat and mouse cell lines derived from kidney, intestine, liver and bone, by immunoprecipitation (using the anti-VDR 9A7 rat monoclonal antibody; Affinity Bioreagents Inc.,

Golden, Colorado) followed by Western blot analysis. The 1d- and 1c-exonspecific antibodies detected the same band in all immunoprecipitations.

DISCUSSION

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The present inventors have identified 5' variant transcripts of the hVDR that suggest the existence of alternative promoters. These transcripts may not have been discriminated in previous Northern analyses because of their similarity in size. Transcription initiation from exons 1a or 1f and alternative splicing generate VDR transcripts that vary in their 5' UTRs but encode the same 427-aa protein. Transcription initiation from exon 1d and alternative splicing generate hVDR transcripts with the potential to encode variant proteins with an additional 50 or 23 aa at the N terminus. There was no evidence that these 5' variants are associated with differences at the 3' end of the transcript. Although isoforms are common in other members of the nuclear receptor superfamily, the only evidence for isoforms of the hVDR is a common polymorphism in the triplet encoding the initiating methionine of the 427-aa form of the VDR that results in initiation of translation at an alternative start codon beginning at the 10th nucleotide down-stream, encoding a protein truncated by 3 aa at the N terminus (5). Similarly, two forms of the avian VDR, differing in size by 14 aa, are generated from a single transcript by alternative translation initiation (6), and in the rat a dominantnegative VDR is generated by intron retention (7).

Heterogeneity in the 5' region is a common feature of other nuclear receptor genes. Tissue-specific alternative-promoter usage generates multiple transcripts of the human estrogen receptor a (ERa), the human and rat mineralocorticoid receptors, and the mouse glucocorticoid receptor (GR), which differ in their 5' UTRs but code for identical proteins. However, other members of the nuclear receptor superfamily have multiple, functionally distinct isoforms arising from differential promoter usage and/or alternative splicing. The generation of N-terminal variant protein isoforms has been described for the progesterone receptor (PR), peroxisome proliferator-activated receptor (PPAR_o), and the retinoid and thyroid receptors. Some receptor isoforms exhibit differential promoter-specific transactivation activity. The N-terminal A/B regions of many nuclear receptor proteins possess a ligand-independent transactivation function (AF1). An AF1

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domain has been demonstrated for the thyroid receptor b1 (TRb1), ER. GR. PR, PPARg, and the retinoid receptors. The activity of the AF1 domain has been shown to vary in both a tissue- and promoter-specific manner. The Nterminal A/B region of nuclear receptors is the least-conserved domain across the family and between receptor subtypes, varying considerably both in length and sequence. The VDR, however, is unusual as its N-terminal A/B region is much shorter than that of other nuclear receptors, with only 23 aa N-terminal to the DNA-binding domain, and deletion of these residues seems to have no effect on VDR function. This region in other receptors is associated with optimal ligand-dependent transactivation and can interact directly with components of the basal transcription complex. Two stretches of basic amino acid residues, RNKKR and RPHRR, in the predicted amino acid sequences of the variant hVDR N termini (Fig. 1C) resemble nuclear localization signals. An N-terminal variant VDR protein therefore might exhibit different transactivation potential, possibly mediated by different protein interactions, or may specify a different subcellular localization. The tissue-specific expression of exon 1f-containing transcripts is mediated by a distal promoter more than 9 kb upstream of exons 1a and 1d. Exon 1f transcripts were detected only in kidney tissue, parathyroid adenoma tissue, and an intestinal cell line, LIM 1863. It is interesting that these tissues represent major target tissues for the calcitropic effects of vitamin D. The absence of 1f-containing transcripts in two other kidney cell lines, HK-2 (proximal tubule) and HEK-293 (embryonal kidney), as well as one other embryonal intestinal cell line, Intestine-407, suggests that the expression of 1f transcripts is cell type-specific. The cell line-specific activity of exon 1f flanking sequences in promoter reporter assays may reflect a requirement for tissue- or cell-specific protein factors to mediate expression from this promoter.

This study has demonstrated that expression of the human VDR gene, which spans more than 60 kb and consists of 14 exons. is under complex transcriptional control by multiple promoters. The expression of multiple exon 1f transcripts is mediated by utilization of a distal tissue-specific promoter. Transcription from a proximal promoter, or promoters, generates multiple variant hVDR transcripts, two of which code for N-terminal variant proteins. Multiple, functionally distinct isoforms mediate the tissue- and/or developmental-specific effects of many members of the nuclear receptor

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superfamily. Although the actual relative abundance of the various transcripts and their levels of translation *in vivo* have not yet been characterized, the results suggest that major variant isoforms of the hVDR exist. Differential regulation of these hVDR gene promoters and of alternative splicing of variant VDR transcripts may have implications for understanding the various actions of 1,25-(OH)₂D₃ in different cell types, and variant VDR transcripts may play a role in tissue specific VDR actions in bone and calcium homeostasis.

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It will be appreciated by persons skilled in the art that numerous variations and/or modifications may be made to the invention as shown in the specific embodiments without departing from the spirit or scope of the invention as broadly described. The present embodiments are, therefore, to be considered in all respects as illustrative and not restrictive.

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Ile	Arg	Cys	Arg 420	His	Pro	Pro	Pro	Gly 425	Ser	His	Leu	Leu	Tyr 430	Ala	Lys
Met	Ile	Gln 435	Lys	Leu	Ala	Asp	Leu 440	Arg	Ser	Leu	Asn	Glu 445	Glu	His	Sei
Lys	Gln 450	Туг	Arg	Cys	Leu	Ser 455	Phe	Gln	Pro	Glu	Cys 460	Ser	Met	Lys	Lev
Thr 465	Pro	Leu	Val	Leu	Glu 470	Val	Phe	Gly	Asn	Glu 475	Ile	Ser			

SEQ ID NO: 10 <211> 450 <212> PRT <213> Homo sapiens

<400> 10

Met Glu Trp Arg Asn Lys Lys Arg Ser Asp Trp Leu Ser Met Val Leu 1 5 10 15

Arg Thr Ala Gly Val Glu Gly Met Glu Ala Met Ala Ala Ser Thr Ser 20 25 30

Leu Pro Asp Pro Gly Asp Phe Asp Arg Asn Val Pro Arg Ile Cys Gly
35 40 45

Val Cys Gly Asp Arg Ala Thr Gly Phe His Phe Asn Ala Met Thr Cys 50 55 60

Glu Gly Cys Lys Gly Phe Phe Arg Arg Ser Met Lys Arg Lys Ala Leu 65 70 75 80

Phe Thr Cys Pro Phe Asn Gly Asp Cys Arg Ile Thr Lys Asp Asn Arg 85 90 95

Arg His Cys Gln Ala Cys Arg Leu Lys Arg Cys Val Asp Ile Gly Met 100 105 110

Met Lys Glu Phe Ile Leu Thr Asp Glu Glu Val Gln Arg Lys Arg Glu 115 120 125

Met Ile Leu Lys Arg Lys Glu Glu Glu Ala Leu Lys Asp Ser Leu Arg 130 135

Pro Lys Leu Ser Glu Glu Gln Gln Arg Ile Ile Ala Ile Leu Leu Asp 145 150 155 160

Ala His His Lys Thr Tyr Asp Pro Thr Tyr Ser Asp Phe Cys Gln Phe 165 170 175

Arg Pro Pro Val Arg Val Asn Asp Gly Gly Gly Ser His Pro Ser Arg 180 185 190

Pro Asn Ser Arg His Thr Pro Ser Phe Ser Gly Asp Ser Ser Ser Ser 195 200 205

Cys Ser Asp His Cys Ile Thr Ser Ser Asp Met Met Asp Ser Ser Ser 210 220

Phe Ser Asn Leu Asp Leu Ser Glu Glu Asp Ser Asp Asp Pro Ser Val 225 230 235

Thr Leu Glu Leu Ser Gln Leu Ser Met Leu Pro His Leu Ala Asp Leu 245 250 255

Val Ser Tyr Ser Ile Gln Lys Val Ile Gly Phe Ala Lys Met Ile Pro 260 265 270

Gly Phe Arg Asp Leu Thr Ser Glu Asp Gln Ile Val Leu Leu Lys Ser 275 280 285

Ser Ala Ile Glu Val Ile Met Leu Arg Ser Asn Glu Ser Phe Thr Met 290 295 300

Asp Asp Met Ser Trp Thr Cys Gly Asn Gln Asp Tyr Lys Tyr Arg Val 305 310 315 320

Ser Asp Val Thr Lys Ala Gly His Ser Leu Glu Leu Ile Glu Pro Leu 325 330 335

Ile Lys Phe Gln Val Gly Leu Lys Lys Leu Asn Leu His Glu Glu Glu 340 345 350

His Val Leu Leu Met Ala Ile Cys Ile Val Ser Pro Asp Arg Pro Gly 355 360 365

Val Gln Asp Ala Ala Leu Ile Glu Ala Ile Gln Asp Arg Leu Ser Asn 370 375 380

Thr Leu Gln Thr Tyr Ile Arg Cys Arg His Pro Pro Pro Gly Ser His 385 390 395 400

Leu Leu Tyr Ala Lys Met Ile Gln Lys Leu Ala Asp Leu Arg Ser Leu 405 410 415

Asn Glu Glu His Ser Lys Gln Tyr Arg Cys Leu Ser Phe Gln Pro Glu 420 425 430

Cys Ser Met Lys Leu Thr Pro Leu Val Leu Glu Val Phe Gly Asn Glu
435 440 445

Ile Ser 450

SEQ ID No: 11

<211> 72

<212> PRT

<213> Homo sapiens

<400> 11

Met Glu Trp Arg Asn Lys Lys Arg Ser Asp Trp Leu Ser Met Val Leu 1 5 10

Arg Thr Ala Gly Val Glu Gly Met Glu Ala Met Ala Ala Ser Thr Ser 20 25 30

Leu Pro Asp Pro Gly Asp Phe Asp Arg Asn Val Pro Arg Ile Cys Gly 35 40

Val Cys Gly Asp Arg Ala Thr Gly Phe His Phe Asn Ala Met Thr Cys 50 60

Glu Gly Cys Lys Gly Phe Phe Arg

SEQ ID NO: 12 <211> 427 <212> PRT <213> Homo sapiens

-

<400> 12

Met Glu Ala Met Ala Ala Ser Thr Ser Leu Pro Asp Pro Gly Asp Phe $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$

Asp Arg Asn Val Pro Arg Ile Cys Gly Val Cys Gly Asp Arg Ala Thr 20 25 30

Gly Phe His Phe Asn Ala Met Thr Cys Glu Gly Cys Lys Gly Phe Phe 35 40 45

Arg Arg Ser Met Lys Arg Lys Ala Leu Phe Thr Cys Pro Phe Asn Gly 50 55 60

Asp Cys Arg Ile Thr Lys Asp Asn Arg Arg His Cys Gln Ala Cys Arg 65 70 75 80

Leu Lys Arg Cys Val Asp Ile Gly Met Met Lys Glu Phe Ile Leu Thr 85 90 95

Asp Glu Glu Val Gln Arg Lys Arg Glu Met Ile Leu Lys Arg Lys Glu
100 105 110

Glu Glu Ala Leu Lys Asp Ser Leu Arg Pro Lys Leu Ser Glu Glu Gln 115 120 125

Gln Arg Ile Ile Ala Ile Leu Leu Asp Ala His His Lys Thr Tyr Asp 130 135 140

Pro Thr Tyr Ser Asp Phe Cys Gln Phe Arg Pro Pro Val Arg Val Asn 145 150 150

Asp Gly Gly Ser His Pro Ser Arg Pro Asn Ser Arg His Thr Pro 165 170 175

Ser Phe Ser Gly Asp Ser Ser Ser Ser Cys Ser Asp His Cys Ile Thr 180 185 190

Ser Ser Asp Met Met Asp Ser Ser Ser Phe Ser Asn Leu Asp Leu Ser 195 200 205

Glu Glu Asp Ser Asp Asp Pro Ser Val Thr Leu Glu Leu Ser Gln Leu 210 215 220

Ser Met Leu Pro His Leu Ala Asp Leu Val Ser Tyr Ser Ile Gln Lys 225 230 235

Val Ile Gly Phe Ala Lys Met Ile Pro Gly Phe Arg Asp Leu Thr Ser 245 250 255

Glu Asp Gln Ile Val Leu Leu Lys Ser Ser Ala Ile Glu Val Ile Met 260 265 270

Leu Arg Ser Asn Glu Ser Phe Thr Met Asp Asp Met Ser Trp Thr Cys 275 280 285

Gly	Asn 290	Gln	Asp	Tyr	Lys	Tyr 295	Arg	Val	Ser	qaA	Val 300	Thr	Lys	Ala	Gly
His 305	Ser	Leu	Glu	Leu	Ile 310	Glu	Pro	Leu	Ile	Lys 315	Phe	Gln	Val	Gly	Let 320
Lys	Lys	Leu	Asn	Leu 325	His	Glu	Glu	Glu	His 330	Val	Leu	Leu	Met	Ala 335	Ile
Cys	Ile	Val	Ser 340	Pro	Asp	Arg	Pro	Gly 345	Val	Gln	Asp	Ala	Ala 350	Leu	Il∈
Glu	Ala	Ile 355	Gln	Asp	Arg	Leu	Ser 360	Asn	Thr	Leu	Gln	Thr 365	Tyr	Ile	Arg
Cys	Arg 370	His	Pro	Pro	Pro	Gly 375	Ser	His'	Leu	Leu	Tyr 380	Ala	Lys	Met	Ile
Gln 385	Lys	Leu	Ala	Asp	Leu 390	Arg	Ser	Leu	Asn	Glu 3,95	Glu	His	Ser	Lys	Gln 400
Tyr	Arg	Cys	Leu	Ser 405	Phe	Gln	Pro	Glu	Cys 410	Ser	Met	Lys	Leu	Thr 415	Pro
Leu	Val	Leu	Glu 420	Val	Phe	Gly	Asn	Glu 425	Ile	Ser					

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Claims:-

- 1. An isolated polynucleotide molecule encoding a human Vitamin D receptor (hVDR) isoform, said polynucleotide molecule comprising a nucleotide sequence which includes sequence that substantially corresponds or is functionally equivalent to that of exon 1d of the human VDR gene.
- 2. A polynucleotide molecule according to claim 1, wherein said nucleotide sequence further includes sequence that substantially corresponds or is functionally equivalent to that of exon 1b and/or exon 1c.
- 3. A polynucleotide molecule according to claim 1, wherein the nucleotide sequence includes:
- (i) sequence that substantially corresponds or is functionally equivalent to that of exons 1d, 1c and 2-9 and encodes a VDR isoform of approximately 477 amino acids,
- (ii) sequence that substantially corresponds or is functionally equivalent to that of exons 1d and 2-9 and encodes a VDR isoform of approximately 450 amino acids, or
- (iii) sequence that substantially corresponds or is functionally equivalent to that of exons 1d and 2-9 and further includes a 152bp intronic sequence and encodes a truncated VDR isoform of approximately 72 amino acids.
- 4. A polynucleotide molecule according to claim 1, wherein the nucleotide sequence substantially corresponds to that shown as SEQ ID NO: 2, SEQ ID NO: 3 or SEQ ID NO: 4.
 - 5. An isolated polynucleotide molecule encoding a human Vitamin D receptor (hVDR), said polynucleotide molecule comprising a nucleotide sequence which includes sequence that substantially corresponds or is functionally equivalent to that of exon 1f and/or 1e of the human VDR gene.
- 6. A polynucleotide molecule according to claim 5, wherein the nucleotide sequence further includes sequence that substantially corresponds or is functionally equivalent to that of exon 1c.

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- 7. A polynucleotide molecule according to claim 5, wherein the nucleotide sequence includes sequence that substantially corresponds or is functionally equivalent to that of exons 1f and 2-9.
- 8. A polynucleotide molecule according to claim 5, wherein the nucleotide sequence substantially corresponds to that shown as SEQ ID NO: 7.
- 9. A plasmid or expression vector including a polynucleotide molecule according to any one of the preceding claims.
 - 10. A host cell transformed with a polynucleotide molecule according to any one of claims 1-8 or a plasmid or expression vector according to claim 9.
 - 11. A host cell according to claim 10, wherein the cell is a mammalian cell.
 - 12. A host cell according to claim 10, wherein the cell is a NIH 3T3 or COS 7 cell.
 - 13. A method of producing a VDR or VDR isoform or functionally equivalent fragments thereof, comprising culturing a host cell of any one of claims 10-12 under conditions enabling the expression of the polynucleotide molecule and, optionally, recovering the VDR or VDR isoform or functionally equivalent fragments thereof.
 - 14. A method according to claim 13, wherein the VDR or VDR isoform or functionally equivalent fragments thereof are expressed onto the host cell membrane or other sub-cellular compartment.
 - 15. A human Vitamin D receptor (hVDR) isoform or functionally equivalent fragment thereof encoded by a polynucleotide molecule according to any one of claims 1-4, said hVDR isoform or functionally equivalent fragment thereof being in a substantially pure form.

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- 16. An antibody or antibody fragment capable of specifically binding to a VDR isoform according to claim 15.
- 17. A non-human animal transformed with a polynucleotide molecule according to any one of claims 1-8.
- 18. A method for detecting agonist and/or antagonist compounds of a VDR isoform of claim 15, comprising contacting said VDR isoform. functionally equivalent fragment thereof or a cell transformed with and expressing a polynucleotide molecule according to any one of claims 1-4, with a test compound under conditions enabling the activation of the VDR isoform or functionally equivalent fragment thereof, and detecting an increase or decrease in the activity of the VDR isoform or functionally equivalent fragment thereof.
- 19. An oligonucleotide or polynucleotide probe comprising a nucleotide sequence of 10 or more nucleotides, the probe comprising a nucleotide sequence such that the probe specifically hybridises to a polynucleotide molecule according to any one of claims 1-8 under high stringency conditions.
- 20. An antisense polynucleotide molecule comprising a nucleotide sequence capable of specifically hybridising to a mRNA molecule which encodes a VDR or VDR isoform encoded by a polynucleotide molecule according to any one of claims 1-8, so as to prevent translation of the mRNA molecule.
- 21. An isolated polynucleotide molecule comprising a nucleotide sequence showing greater than 75% sequence identity to:
- (i) 5'TGCGACCTTGGCGGTGAGCCTGGGGACAGGGGTGAGGCCAGAGA CGGACGGACGCAGGGCCCCGGCCCAAGGCGAGGAAACAGCGGCACTA AGGCAGAAAGGAAGAGGGCGGTGTTTCACCCGCAGCCCAATCCATCAC TCAGCAACTCCTAGACGCTGGTAGAAAGTTCCTCCGAGGAGCCTGCCATC CAGTCGTGCGTGCAG3' (SEQ ID NO: 5)

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- (ii) 5'AGGCAGCATGAAACAGTGGGATGTGCAGAGAGAAGATCTGGGTC CAGTAGCTCTGACACTCCTCAGCTGTAGAAACCTTGACAACTCTGCACAT CAGTTGTACAATGGAACGGTATTTTTTACTCTTCATGTCTGAAAAAGGCTA TGATAAAGATCAA3' (SEQ ID NO: 6), or
- 22. An isolated polynucleotide molecule comprising a nucleotide sequence showing greater than 85% sequence identity to:
- (ii) 5'AGGCAGCATGAAACAGTGGGATGTGCAGAGAGAGATCTGGGTC

 CAGTAGCTCTGACACTCCTCAGCTGTAGAAACCTTGACAACTCTGCACAT

 CAGTTGTACAATGGAACGGTATTTTTTACTCTTCATGTCTGAAAAGGCTA

 TGATAAAGATCAA3' (SEQ ID NO: 6), or
- (iii) 5'GTTTCCTTCTGTCGGGGCGCCCTTGGCATGGAGTGGAGGAATA
 AGAAAAGGAGCGATTGGCTGTCGATGGTGCTCAGAACTGCTGGAGTGGA
 GG3' (SEQ ID NO: 1).
 - 23. An isolated polynucleotide molecule comprising a nucleotide sequence showing greater than 95% sequence identity to:

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- (ii) 5'AGGCAGCATGAAACAGTGGGATGTGCAGAGAGAAGATCTGGGTC CAGTAGCTCTGACACTCCTCAGCTGTAGAAACCTTGACAACTCTGCACAT CAGTTGTACAATGGAACGGTATTTTTTACTCTTCATGTCTGAAAAAGGCTA 'TGATAAAGATCAA3' (SEQ ID NO: 6), or
- (iii) 5'GTTTCCTTCTGTCGGGGCGCCCTTGGCATGGAGTGGAGGAATA AGAAAAGGAGCGAT'I'GGCTGTCGATGGTGCTCAGAACTGCTGGAGTGGA GG3' (SEQ ID NO: 1)
- 24. An isolated polynucleotide molecule comprising nucleotide sequence substantially corresponding to:
 - (i) 5'TGCGACCTTGGCGGTGAGCCTGGGGACAGGGGTGAGGCCAGAGA CGGACGGACGCAGGGGCCCCAAGGCGAGGAGAACAGCGGCACTA AGGCAGAAAGGAAGAGGGCGGTGTGTTCACCCGCAGCCCAATCCATCAC TCAGCAACTCCTAGACGCTGGTAGAAAGTTCCTCCGAGGAGCCTGCCATC CAGTCGTGCGTGCAG3' (SEQ ID NO: 5)
 - (ii) 5'AGGCAGCATGAAACAGTGGGATGTGCAGAGAGAAGATCTGGGTC CAGTAGCTCTGACACTCCTCAGCTGTAGAAACCTTGACAACTCTGCACAT CAGTTGTACAATGGAACGGTATTTTTTACTCTTCATGTCTGAAAAGGCTA TGATAAAGATCAA3' (SEQ ID NO: 6), or
- (iii) 5'GTTTCCTTCTGTCGGGGCGCCCTTGGCATGGAGTGGAGGAATA
 AGAAAAGGAGCGATTGGCTGTCGATGGTGCTCAGAACTGCTGGAGTGGA
 GG3' (SEQ ID NO: 1)

Transcript 9:

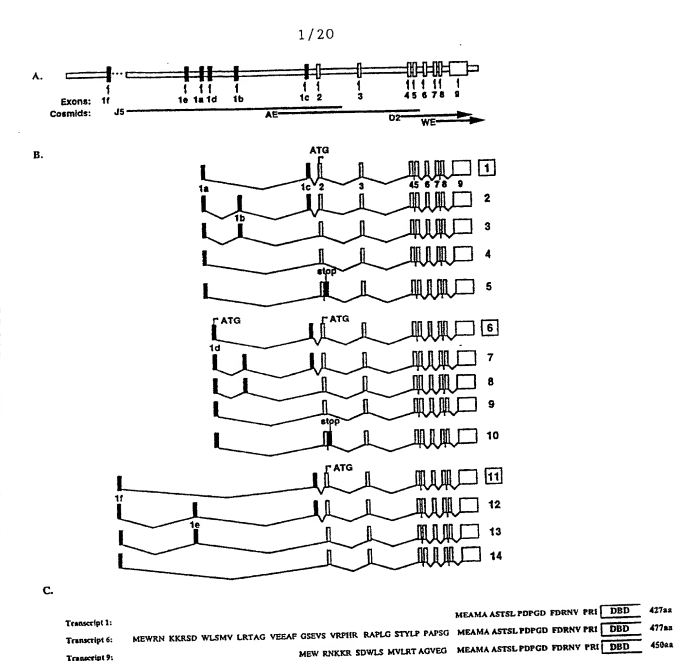


FIGURE 1

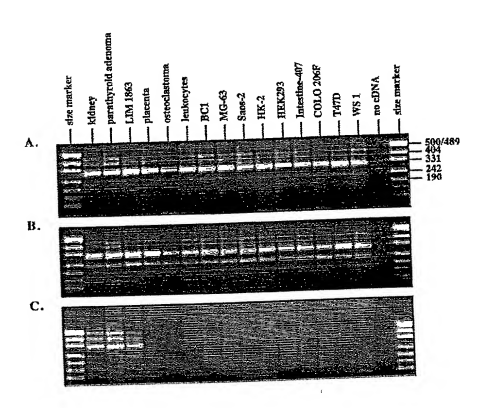


FIGURE 2

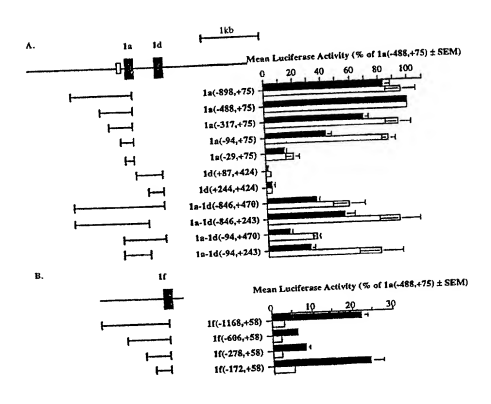


FIGURE 3

- C. 5'...tgtttttagAGGCAGCATGAAACAGTGGGATGTGCAGAGAGAAGATCTGGGTCCAGTAGCTCTGACACTCCTCAGCTGTAGAAACCTTGACAACTCTGCACATCAGTTGTACAATGGAACGGTATTTTTTACTCTTCATGTCTGAAAAAGGCTATGATAAAAGATCAAgtaagatatt...3'

FIGURE 5	TR	ANS	CRIPT	6						
(Sequence F	ange: 1 t	0 14	63)							
	*	10		20		30 *	4	40		50
	GTTTCCTT	• •	ירינויטיניר <i>י</i> נ "		CGCCmu		ጥርርኔርጥ	-	GAATAA	C
	CAAAGGAA									
							MetGluT			
		60		70		80		90		100
	*	*	*	*	*	*	*	*	*	*
	AGGAGCGA TCCTCGCT									
	ArgSerAs									
	J						•			
	_	10		120		130		140		150
	*	*	*	*	*	*	*	*	*	*
	CTTTGGGT GAAACCCA									
	PheGlyS									
							_	-		_
	1	60		170		180		190		200
	* CCACTTAC	* ~~ ~	*	*	*	*	# ###	ייי א אייי	eccece	* CAGC
	GGTGAATG									
	SerThrTyr									
	2	10		220		230		240		250
	*	*	*	*	*	*	*	*	*	*
	ACTTCCCT									
	TGAAGGGA ThrSerLe	CG G	ACTGG	GACC	TCTGAA	ACTG	GCCTTG	CACG	GGGCCT	AGAC 1aCve>
	Thrserie	u PI	OASDP	rogry	y Aspri	iewsb	ALGASI	val	TONIGI	recyse
	2	60		270		280		290		300
	*	*	*	*	*	*	*	*	*	*
	TGGGGTGI									
	ACCCCACA GlyValC									
	Giyvaic	ys c	TAYSD.	Arg A	ALATHIC	at A Liv	e nisen	CASII	Aranec	
	3	10		320		330		340		350
	*	*	*	*	*	*	*	*	*	*
	GTGAAGGC	TG (CAAAGG	CTTC	TTCAGO	CGAA	GCATGA	AGCG	GAAGGC	ACTA MGMT
	CACTTCCC CysGluGly	Cys	LysG1	yPhe	PheArg	JArg	SerMetI	ysAr	g LysAl	aLeu>
	3	60		370		380		390	•	400
	*	*	*	*	*	*	*	*	*	*
	TTCACCTO	CC (CCTTCA	ACGG	GGACT	GCCGC	ATCACO	ZAAGG	ACAACO	CGACG
	AAGTGGAG PheThrCy	.GG (rs Pi	sGAAGT roPhe≥	rucc snG1	CCTGAC Odea v	vsAra	IleTh	Lys	Aggaga Aggaga	rgArg>
					,		•	-	-	

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	0,	/ 20		
410	420	430	: 440	450
* *	* *	* *	* *	* *
	GCCTGCCGGC			
	CGGACGGCCG			
HISCYSGIN	AlaCysArg I	.euLysArgCys	s valaspile	Giymetmet>
460	470	480	490	500
* *	* *	* *	* *	* *
	TCTGACAGAT			
	AGACTGTCTA			
LysGluPheIle	e LeuThrAsp	GluGluVal C	InArgLysAr	g GluMetIle>
		~~~	540	
510 * *	520 * *	530 * *	540 * *	550 * *
	AGGAGGAGGA			
	TCCTCCTCCT			
				ArgProLysLeu>
neanysary :	nyseruerueru	. WIGHERHAS	Aspect Bed ?	argriony sheu/
560	570	580	590	600
* *	* *	* *	* *	* *
GTCTGAGGAG	CAGCAGCGCA	TCATTGCCAT	ACTGCTGGAC	GCCCACCATA
CAGACTCCTC	GTCGTCGCGT	AGTAACGGTA	TGACGACCTG	CGGGTGGTAT
SerGluGlu	GlnGlnArg	TleIleAlaIle	e LeuLeuAsp	AlaHisHis>
610	620	630	640	650
* *	* *	* *	* *	* *
	CCCCACCTAC			
	GGGGTGGATG			
Lys'inr'iyrAs	p Prolnriyr	Seraspene (	LysGinPhear	g ProProVal>
660	670	680	690	700
* *	* *	* *	* *	* *
CGTGTGAATG	ATGGTGGAGG	GAGCCATCCT	TCCAGGCCCA	ACTCCAGACA
				TGAGGTCTGT
				AsnSerArgHis>
_				
710	720	730	740	750
* *	* *	* *	* *	* *
CACTCCCAGC	TTCTCTGGGG	ACTCCTCCTC	CTCCTGCTCA	GATCACTGTA
GTGAGGGTCG	AAGAGACCCC	TGAGGAGGAG	GAGGACGAGT	CTAGTGACAT
ThrProSer	PheSerGly	AspSerSerse:	r sercysser	AspHisCys>
760	770	780	790	800
* *	* *	* *	* *	· -
<b>ጥር ልርርጥርጥጥር</b>	AGACATGATG	GACTCGTCCA	GCTTCTCCAA	TCTGGATCTG
AGTGGAGAAG	TCTGTACTAC	CTGAGCAGGT	CGAAGAGGTT	AGACCTAGAC
IleThrSerSe	r AspMetMet	AspSerSer	SerPheSerAs	n LeuAspLeu>
810				
* *				
AGTGAAGAAG	ATTCAGATGA	CCCTTCTGTG	ACCCTAGAGC	TGTCCCAGCT
TCACTTCTTC	TAAGTCTACT	GGGAAGACAC	TGGGATCTCG	ACAGGGTCGA
SerGluGlu	AspserAspAs	p Proserval	THIPERGIR	LeuSerGlnLeu>

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	7	/20		
860 * *	870 * *	880 * *	. 890 * *	900
		CTGACCTGGT		
		GACTGGACCA		
SerMetLeu	ProHisLeu A	\laAspLeuVal	l SerTyrSer	IleGlnLys>
910	920	930	940	950
* *	* *	* *	* *	* *
TCATTGGCTT	TGCTAAGATG	ATACCAGGAT	TCAGAGACCT	CACCTCTGAG
AGTAACCGAA	ACGATTCTAC	TATGGTCCTA	AGTCTCTGGA	GTGGAGACTC
VallleGlyPhe				
-	-	-		
960	970	980	990	1000
* *	* *	* *	* *	* *
GACCAGATCG	TACTGCTGAA	GTCAAGTGCC	ATTGAGGTCA	TCATGTTGCG
CTGGTCTAGC	ATGACGACTT	CAGTTCACGG	TAACTCCAGT	AGTACAACGC
				leMetLeuArg>
vabaturie (	arneaneanys	berberara	IICGIUVAI .	remetheurig>
1010	1020	1030	1040	1050
* *	* *	* *	* *	* *
CTCCAATGAG	TOCTTOACCA	TGGACGACAT	GTCCTGGACC	ጥርጥርርርር እ እ ር ር
		ACCTGCTGTA		
		MetAspAspMet		
pervendia	perment t	е сивриврие (	. Dellipini	CysGlyRsii>
1060	1070	1080	1090	1100
* *	* *	* *	* *	* *
AAGACTACAA	GTACCGCGTC	AGTGACGTGA	CCAAAGCCGG	ACACAGCCTG
TTCTGATGTT	CATGGCGCAG	TCACTGCACT	GGTTTCGGCC	TGTGTCGGAC
GlnAspTyrLys	TyrArgVal	SerAspVal T	hrLysAlaGly	HisSerLeu>
1110	1120	1130	1140	1150
* *	* *	* *	* *	* *
		CAAGTTCCAG		
CTCGACTAAC	TCGGGGAGTA	GTTCAAGGTC	CACCCTGACT	TCTTCGACTT
GluLeuIle (	GluProLeuIle	LysPheGln	ValGlyLeu I	ysLysLeuAsn>
1160	1170	1180	1190	1200
* *	* *	* *	* *	* *
CTTGCATGAG	GAGGAGCATG	TCCTGCTCAT	GGCCATCTGC	ATCGTCTCCC
GAACGTACTC	CTCCTCGTAC	AGGACGAGTA	CCGGTAGACG	TAGCAGAGGG
LeuHisGlu	GluGluHis V	/alLeuLeuMet	AlaIleCys	<pre>IleValSer&gt;</pre>
1210				1250
* *				
CAGATCGTCC	TGGGGTGCAG	GACGCCGCGC	TGATTGAGGC	CATCCAGGAC
				GTAGGTCCTG
ProAspArgPro	o GlyValGln	AspAlaAla	LeulleGluAla	a IleGlnAsp>
1260		1280		1300
* *	* *	* *	* *	* *
				ACCCGCCCC
				TGGGCGGGG
ArgLeuSer .	AsnThrLeuGl:	n ThrTyrIle	ArgCysArg	HisProProPro;

8/20 1330 1340 1320 1350 GGGCAGCCAC CTGCTCTATG CCAAGATGAT CCAGAAGCTA GCCGACCTGC CCCGTCGGTG GACGAGATAC GGTTCTACTA GGTCTTCGAT CGGCTGGACG GlySerHis LeuLeuTyr AlaLysMetIle GlnLysLeu AlaAspLeu> 1380 1390 1400 GCAGCCTCAA TGAGGAGCAC TCCAAGCAGT ACCGCTGCCT CTCCTTCCAG CGTCGGAGTT ACTCCTCGTG AGGTTCGTCA TGGCGACGGA GAGGAAGGTC ArgSerLeuAsn GluGluHis SerLysGln TyrArgCysLeu SerPheGln> 1430 1450 CCTGAGTGCA GCATGAAGCT AACGCCCCTT GTGCTCGAAG TGTTTGGCAA GGACTCACGT CGTACTTCGA TTGCGGGGAA CACGAGCTTC ACAAACCGTT

ProGluCys SerMetLysLeu ThrProLeu ValLeuGlu ValPheGlyAsn>

1460

* *
TGAGATCTCC TGA
ACTCTAGAGG ACT
GluIleSer ***>

FIGURE 6	TRANS	CRIPT 9			
(Sequence Ra	inge: 1 to	1382)			
	10	20	30	40	50
	* *	* *	* *	* *	* *
	GTTTCCTTCT			TGGAGTGGAG ACCTCACCTC	
	CZMMICOJIMICI.		1	MetGluTrpArg	AsnLysLys>
	60 * *	70 * *	80 * *	90 * *	100 * *
	AGGAGCGATT	GGCTGTCGAT	GGTGCTCAGA	ACTGCTGGAG	TGGAGGGGAT
	TCCTCGCTAA	CCGACAGCTA	CCACGAGTCT	TGACGACCTC	ACCTCCCCTA
	ArgSerAsp	TrpLeuSerMe	t ValLeuArg	ThrAlaGly V	/alGluGlyMet>
	110	120	130	140	150
	* *	* *	* *	* *	* *
	GGAGGCAATG	GCGGCCAGCA	CTTCCCTGCC	TGACCCTGGA	GACTTTGACC
	CCTCCGTTAC	CGCCGGTCGT	GAAGGGACGG	ACTGGGACCT	CTGAAACTGG
	GluAlaMet	AlaAlaSer	ThrSerLeuPr	o AspProGly	AspPheAsp>
	160 * *	170 * *	180 * *	190 * *	200 * *
	GGAACGTGCC	CCGGATCTGT	GGGGTGTGTG	GAGACCGAGC	CACTGGCTTT
	CCTTGCACGG	GGCCTAGACA	CCCCACACAC	CTCTGGCTCG	GTGACCGAAA
	ArgAsnValPr	o ArgIleCys	GlyValCys	GlyAspArgAla	a ThrGlyPhe>
	210	220	230	240	250
	* *				* *
	GTGAAGTTAC	GATACTGGAC	ACTTCCGACG	AAAGGCTTCT TTTCCGAAGA	AGTCCGCTTC
	HisPheAsn	AlaMetThrCy	s GluGlyCys	LysGlyPhe	PheArgArgSer>
	260		280	290	300 * *
	CATGAAGCG	AAGGCACTAT	TCACCTGCCC	CTTCAACGGG	GACTGCCGCA
	GTACTTCGC	TTCCGTGAT	AGTGGACGG	GAAGTTGCCC	CTGACGGCGT
	MetLysArg	J LysAlaLeu	PheThrCysPi	o PheAsnGly	AspCysArg>
	310	320	330	340	350
	*	* * :	* * '	* * *	* *
	TCACCAAGG	A CAACCGACG	C CACTGCCAG	CCTGCCGGC1	CAAACGCTGT
	AGTGGTTCC	r GTTGGCTGC	G GTGACGGTC	C GGACGGCCGA	GTTTGCGACA
	IleThrLysA	sp AsnArgAr	g HisCysGln	AlaCysArgLe	eu LysArgCys>
		* *	* *	* * 1	k * *
	GTGGACATC	G GCATGATGA	A GGAGTTCAT	T CTGACAGAT	AGGAAGTGCA
	CACCTGTAG	C CGTACTACT	T CCTCAAGTA	A GACTGTCTAC	C TCCTTCACGT
	ValAspIle	GlyMetMetL	ys GluPheIl	e LeuThrAsp	GluGluValGln>

	,			
410	420 * *	430 * *	· 440	450 * *
GAGGAAGCGG	GAGATGATCC	TGAAGCGGAA	GGAGGAGGAG	GCCTTGAAGG
CTCCTTCGCC	CTCTACTAGG	ACTTCGCCTT	CCTCCTCCTC	CGGAACTTCC
ArgLysArg	GluMetIle I	euLysArg <b>L</b> y:	s GluGluGlu	AlaLeuLys>
460 * *	470 * *	480 * *	490 * *	500 * *
ACAGTCTGCG	GCCCAAGCTG	TCTGAGGAGC	AGCAGCGCAT	CATTGCCATA
TGTCAGACGC	CGGGTTCGAC	AGACTCCTCG	TCGTCGCGTA	GTAACGGTAT
AspSerLeuAr	r Prolysleu	SerGluGlu (	GlnGlnArgIle	e IleAlaIle>
napoci nemi.	g		_	
510	520	530	540	550
* *	* *	* *	* *	* *
CTGCTGGACG	CCCACCATAA	GACCTACGAC	CCCACCTACT	CCGACTTCTG
GACGACCTGC	GGGTGGTATT	CTGGATGCTG	GGGTGGATGA	GGCTGAAGAC
LeuLeuAsp .	AlaHisHisLy	s ThrTyrAsp	ProThrTyr	SerAspPheCys>
560	570	580	590	600
* *	* *	* *	* *	* *
CCAGTTCCGG	CCTCCAGTTC	GTGTGAATGA	TGGTGGAGGG	AGCCATCCTT
GGTCAAGGCC	GGAGGTCAAG	CACACTTACT	ACCACCTCCC	TCGGTAGGAA
GlnPheArg	ProProVal	ArgValAsnAs	p GlyGlyGly	SerHisPro>
610	620	630	640	650
* *		* *	* *	* *
CCAGGCCCAA	CTCCAGACAC	ACTCCCAGCT	TCTCTGGGGA	CTCCTCCTCC
CCTCCCCCTT	GAGGTCTGTG	TGAGGGTCGA	. AGAGACCCCT	GAGGAGGAGG
SerArgProAs	n SerArgHis	ThrProSer	PheSerGlyAs	p SerSerSer>
660	670	680	690	700
* *	* * *			
TCCTGCTCAG	ATCACTGTAT	CACCTCTTCA	GACATGATGG	ACTCGTCCAG
AGGACGAGTO	TAGTGACATA	GTGGAGAAGI	CIGIACIACC	NCorcorcer>
SerCysSer	AspHisCysIl	e ThrSerSer	AspMetMet	AspSerSerSer>
		. 720	740	750
710				
	t * *		•	
CTTCTCCAA	r CTGGATCTGA	GIGAAGAAGA	TICAGAIGAC	CCTTCTGTGA
GAAGAGGTT	A GACCTAGACT	CACTICITO	AAGTCTACTC	GGAAGACACT
PheSerAsı	n LeuAspLeu	SerGluGluAs		ProSerVal>
76	-			) 800 * * *
			* * *	• •
CCCTAGAGC	T GTCCCAGCT	C TCCATGCTG	C CCCACCTGG	TGACCTGGTC
CCCATCTC	A CAGGGTCGAG	G AGGTACGAC	G GGGTGGACC	3 ACTGGACCAG
ThrLeuGluL	eu SerGlnLe	u SerMetLeu	ProHisLeuA	la AspLeuVal>
81				
*			••	• •
AGTTACAGC	A TCCAAAAGG	T CATTGGCTT	T GCTAAGATG	A TACCAGGATT
ጥር እ አጥርጥር ር	T AGGTTTTCC	A GTAACCGAA	A CGATTCTAC	T ATGGTCCTAA
SerTyrSer	IleGlnLysV	al IleGlyPh	e AlaLysMet	IleProGlyPhe

860 * *	870 * *	880 * *	. 890 * *	900
	ACCTCTGAGG			
	TGGAGACTCC			
	ThrSerGlu A			
5		•	_	
910	920	930	940	950
* *	* *	* *	* *	* *
	CATGTTGCGC			
	GTACAACGCG			
TieGiuvaille	e metheuarg	Serasngiu s	errhernime	t AspAspMet>
960	970	980	990	1000
* *	* *	* *	* *	* *
TCCTGGACCT	GTGGCAACCA	AGACTACAAG	TACCGCGTCA	GTGACGTGAC
	CACCGTTGGT			
SerTrpThr (	CysGlyAsnGlr	AspTyrLys	TyrArgVal :	SerAspValThr>
		4000	1010	1050
1010	1020	1030	1040 * * *	1050 * *
	CACAGCCTGG			
	GTGTCGGACC			
LvsAlaGlv	HisSerLeu (	SluLeuIleGlu	ProLeuIle	LysPheGln>
_				
1060	1070	1080	1090	1100
* *	* *	* *	* *	* *
TGGGACTGAA	GAAGCTGAAC CTTCGACTTG	TTGCATGAGG	AGGAGCATGT	CCTGCTCATG
WalGlyLeuly	CITCGACTIG	LeuHisGlu (	RluGluHisVa	l LeuLeuMet>
vardrybeaby.	a Tybecamon	204112024		
1110	1120	1130	1140	1150
* *	* *	* *	* *	* *
GCCATCTGCA	TCGTCTCCCC	AGATCGTCCT	GGGGTGCAGG	ACGCCGCGCT
CGGTAGACGT	AGCAGAGGGG	TCTAGCAGGA	CCCCACGTCC	TGCGGCGCGA
AlaIleCys	IleValSerPro	o Aspargero,	GiyvaiGin .	AspAlaAlaLeu>
1160	. 1170	1180	1190	1200
* *	* *	* *	* *	* *
GATTGAGGCC	ATCCAGGACC	GCCTGTCCAA	CACACTGCAG	ACGTACATCC
CTAACTCCGG	TAGGTCCTGG	CGGACAGGTT	GTGTGACGTC	TGCATGTAGG
IleGluAla	IleGlnAsp	ArgLeuSerAs:	n ThrLeuGln	ThrTyrIle>
	1000	1000	1040	1250
1210		1230		
				CAAGATGATC
CGACGGCGGT	GGGCGGGGGC	CCGTCGGTGG	ACGAGATACG	GTTCTACTAG
ArgCysArgHi	s ProProPro	GlySerHis	LeuLeuTyrAl	a LysMetIle>
1260				
* *				
CAGAAGCTAG	CCGACCTGCG	CAGCCTCAAT  CTCCCACTTA	CTCCTCCTC1	CCAAGCAGTA GGTTCGTCAT
Glulvsleu	AlaAspLeuAr	g SerLeuAsn	GluGluHis	SerLysGlnTyr>
		-		•

CCGCTGCCTC TCCTTCCAGC CTGAGTGCAG CATGAAGCTA ACGCCCCTTG GGCGACGGAG AGGAAGGTCG GACTCACGTC GTACTTCGAT TGCGGGGAAC ArgCysLeu SerPheGln ProGluCysSer MetLysLeu ThrProLeu>

1360 1370 1380

TGCTCGAAGT GTTTGGCAAT GAGATCTCCT GA ACGAGCTTCA CAAACCGTTA CTCTAGAGGA CT ValLeuGluVal PheGlyAsn GlulleSer ***> FIGURE 7 TRANSCRIPT 10

(Sequence Range: 1 to 1534)

10 20 30 40 50 MetGluTrpArg AsnLysLys> 60 70 80 90 100 AGGAGCGATT GGCTGTCGAT GGTGCTCAGA ACTGCTGGAG TGGAGGGGAT TCCTCGCTAA CCGACAGCTA CCACGAGTCT TGACGACCTC ACCTCCCCTA ArgSerAsp TrpLeuSerMet ValLeuArg ThrAlaGly ValGluGlyMet> 120 130 140 GGAGGCAATG GCGGCCAGCA CTTCCCTGCC TGACCCTGGA GACTTTGACC CCTCCGTTAC CGCCGGTCGT GAAGGGACGG ACTGGGACCT CTGAAACTGG GluAlaMet AlaAlaSer ThrSerLeuPro AspProGly AspPheAsp> 170 180 190 GGAACGTGCC CCGGATCTGT GGGGTGTGTG GAGACCGAGC CACTGGCTTT CCTTGCACGG GGCCTAGACA CCCCACACAC CTCTGGCTCG GTGACCGAAA ArgAsnValPro ArgIleCys GlyValCys GlyAspArgAla ThrGlyPhe> 220 230 240 CACTTCAATG CTATGACCTG TGAAGGCTGC AAAGGCTTCT TCAGGTGAGC GTGAAGTTAC GATACTGGAC ACTTCCGACG TTTCCGAAGA AGTCCACTCG HisPheAsn AlaMetThrCys GluGlyCys LysGlyPhe PheArg*** 280 290 260 270 * * CCCCTCCCA GGCTCTCCCC AGTGGAAAGG GAGGGAGAAG AAGCAAGGTG GGGGGAGGGT CCGAGAGGGG TCACCTTTCC CTCCCTCTTC TTCGTTCCAC 330 340 320 TTTCCATGAA GGGAGCCCTT GCATTTTTCA CATCTCCTTC CTTACAATGT AAAGGTACTT CCCTCGGGAA CGTAAAAAGT GTAGAGGAAG GAATGTTACA 380 390 400 360 370

CCATGGAACA TGCGGCGCTC ACAGCCACAG GAGCAGGAGG GTCTTGGCGA GGTACCTTGT ACGCCGCGAG TGTCGGTGTC CTCGTCCTCC CAGAACCGCT

	1-	1/20		
410 * *	420 * *		: 440 * *	450 * *
AGCATGAAGC	GGAAGGCACT	ATTCACCTGC		
		TAAGTGGACG		
460	470	480 * *	490 * *	500
* *	* *			* *
		GCCACTGCCA		
		CGGTGACGGT		
* *	520 * *	530 * *	540 * *	* *
GTGTGGACAT	CGGCATGATG	AAGGAGTTCA	TTCTGACAGA	TGAGGAAGTG
CACACCTGTA	GCCGTACTAC	TTCCTCAAGT	AAGACTGTCT	ACTCCTTCAC
560	570	580	590	600
* *	* *	* *	* *	<del>-</del>
CAGAGGAAGC	GGGAGATGAT	CCTGAAGCGG	AAGGAGGAGG	AGGCCTTGAA
		GGACTTCGCC		
610	620	630		650 * *
* *		* *		
		TGTCTGAGGA		
CCTGTCAGAC	GCCGGGTTCG	ACAGACTCCT	CGTCGTCGCG	TAGTAACGGT
660 * *	670 * *	680 * *	690 * *	700 * *
TACTGCTGGA	CGCCCACCAT	AAGACCTACG	ACCCCACCTA	CTCCGACTTC
		TTCTGGATGC		
710	720	730		750
* *	* *	* *	* *	
	GGCCTCCAGT	TCGTGTGAAT	GATGGTGGAG	GGAGCCATCC
	GGCCTCCAGT		GATGGTGGAG	GGAGCCATCC
	GGCCTCCAGT	TCGTGTGAAT	GATGGTGGAG CTACCACCTC 790	GGAGCCATCC CCTCGGTAGG 800
ACGGTCAAGG 760 * *	GGCCTCCAGT CCGGAGGTCA 770 * *	TCGTGTGAAT AGCACACTTA 780 * *	GATGGTGGAG CTACCACCTC 790 * *	GGAGCCATCC CCTCGGTAGG 800 * *
ACGGTCAAGG 760 * * TTCCAGGCCC	GGCCTCCAGT CCGGAGGTCA 770 * *	TCGTGTGAAT AGCACACTTA 780 * * ACACTCCCAG	GATGGTGGAG CTACCACCTC 790 * *	GGAGCCATCC CCTCGGTAGG 800 * *
ACGGTCAAGG 760 * * TTCCAGGCCC	GGCCTCCAGT CCGGAGGTCA 770 * *	TCGTGTGAAT AGCACACTTA 780 * *	GATGGTGGAG CTACCACCTC 790 * *	GGAGCCATCC CCTCGGTAGG 800 * *
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG	TCGTGTGAAT AGCACACTTA 780 * * ACACTCCCAG TGTGAGGGTC	GATGGTGGAG CTACCACCTC 790 * * CTTCTCTGGG GAAGAGACCC	GGAGCCATCC CCTCGGTAGG 800 * * GACTCCTCCT CTGAGGAGGA 850
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 *	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * *	TCGTGTGAAT AGCACACTTA 780 * * ACACTCCCAG TGTGAGGGTC 830 *	GATGGTGGAG CTACCACCTC 790 * * CTTCTCTGGG GAAGAGACCC 840 *	GGAGCCATCC CCTCGGTAGG  * * GACTCCTCCT CTGAGGAGGA  * *
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 * *	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * * AGATCACTGT	TCGTGTGAAT AGCACACTTA  780  * ACACTCCCAG TGTGAGGGTC  830  * ATCACCTCTT	GATGGTGGAG CTACCACCTC 790 * * CTTCTCTGGG GAAGAGACCC 840 * * CAGACATGAT	GGAGCCATCC CCTCGGTAGG   * * GACTCCTCCT CTGAGGAGGA   * * GGACTCGTCC
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 * *	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * * AGATCACTGT	TCGTGTGAAT AGCACACTTA 780 * * ACACTCCCAG TGTGAGGGTC 830 *	GATGGTGGAG CTACCACCTC 790 * * CTTCTCTGGG GAAGAGACCC 840 * * CAGACATGAT	GGAGCCATCC CCTCGGTAGG   * * GACTCCTCCT CTGAGGAGGA   * * GGACTCGTCC
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 * * CCTCCTGCTC GGAGGACGAG	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * * AGATCACTGT TCTAGTGACA	TCGTGTGAAT AGCACACTTA  780  * ACACTCCAG TGTGAGGGTC  830  * ATCACCTCTT TAGTGGAGAA	GATGGTGGAG CTACCACCTC  790  * * CTTCTCTGGG GAAGAGACCC  840  * CAGACATGAT GTCTGTACTA  890	GGAGCCATCC CCTCGGTAGG   800  * * GACTCCTCCT CTGAGGAGGA  850  * GGACTCGTCC CCTGAGCAGG
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 * * CCTCCTGCTC GGAGGACGAG 860 * *	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * * AGATCACTGT TCTAGTGACA 870 * *	TCGTGTGAAT AGCACACTTA  780  * * ACACTCCAG TGTGAGGGTC  830  * ATCACCTCTT TAGTGGAGAA  880  * *	GATGGTGGAG CTACCACCTC  790  * * CTTCTCTGGG GAAGAGACCC  840  * CAGACATGAT GTCTGTACTA  890  * *	GGAGCCATCC CCTCGGTAGG  800  * * GACTCCTCCT CTGAGGAGGA  850  * GGACTCGTCC CCTGAGCAGG  900  *
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 * * CCTCCTGCTC GGAGGACGAG 860 * * AGCTTCTCCA	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * * AGATCACTGT TCTAGTGACA 870 * * ATCTGGATCT	TCGTGTGAAT AGCACACTTA  780  * ACACTCCAG TGTGAGGGTC  830  * ATCACCTCTT TAGTGGAGAA	GATGGTGAG CTACCACCTC  790  * * CTTCTCTGGG GAAGAGACCC  840  * CAGACATGAT GTCTGTACTA  890  * * GATTCAGATG	GGAGCCATCC CCTCGGTAGG   800  * * GACTCCTCCT CTGAGGAGGA  850  * * GGACTCGTCC CCTGAGCAGG  900  * * ACCCTTCTGT
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 * * CCTCCTGCTC GGAGGACGAG 860 * * AGCTTCTCCA	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * * AGATCACTGT TCTAGTGACA 870 * ATCTGGATCT TAGACCTAGA	TCGTGTGAAT AGCACACTTA  780  * * ACACTCCCAG TGTGAGGGTC  830  * * ATCACCTCTT TAGTGGAGAA  880  * GAGTGAAGAA CTCACTTCTT	GATGGTGAG CTACCACCTC  790  * CTTCTCTGGG GAAGAGACCC  840  * CAGACATGAT GTCTGTACTA  890  * GATTCAGATG CTAAGTCTAC	GGAGCCATCC CCTCGGTAGG   * * GACTCCTCT CTGAGGAGGA  * * GGACTCGTCC CCTGAGCAGG  900  * ACCCTTCTGT TGGGAAGACA
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 * * CCTCCTGCTC GGAGGACGAG 860 * * AGCTTCTCCA TCGAAGAGGT	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * * AGATCACTGT TCTAGTGACA 870 * ATCTGGATCT TAGACCTAGA	TCGTGTGAAT AGCACACTTA  780  * * ACACTCCCAG TGTGAGGGTC  830  * * ATCACCTCTT TAGTGGAGAA  880  * GAGTGAAGAA CTCACTTCTT	GATGGTGAG CTACCACCTC  790  * * CTTCTCTGGG GAAGAGACCC  840  * * CAGACATGAT GTCTGTACTA  890  * GATTCAGATG CTAAGTCTAC  940	GGAGCCATCC CCTCGGTAGG   * * GACTCCTCT CTGAGGAGGA  * * GGACTCGTCC CCTGAGCAGG  900  * ACCCTTCTGT TGGGAAGACA
ACGGTCAAGG 760 * * TTCCAGGCCC AAGGTCCGGG 810 * * CCTCCTGCTC GGAGGACGAG 860 * * AGCTTCTCCA TCGAAGAGGT	GGCCTCCAGT CCGGAGGTCA 770 * * AACTCCAGAC TTGAGGTCTG 820 * * AGATCACTGT TCTAGTGACA 870 * ATCTGGATCT TAGACCTAGA 920 * *	TCGTGTGAAT AGCACACTTA  780  * * ACACTCCCAG TGTGAGGGTC  830  * * ATCACCTCTT TAGTGGAGAA  880  * * GAGTGAAGAA CTCACTTCTT  930  * *	GATGGTGAG CTACCACCTC  790  * CTTCTCTGGG GAAGAGACCC  840  * CAGACATGAT GTCTGTACTA  890  * GATTCAGATG CTAAGTCTAC  940  *	GGAGCCATCC CCTCGGTAGG

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960	970 * *	<b>9</b> 80 * *	. 990 * *	1000
			TTGCTAAGAT	
AGTCAATGTC	GTAGGTTTTC	CAGTAACCGA	AACGATTCTA	CTATGGTCCT
4040	4000	1020	1040	1050
1010	1020	1030	1040	1050
* *	* *	* *	* *	* *
TOTAL ACADA	מיכיא ביכישיבישיביא	CCACCACATC	GTACTGCTGA	አርጥር አ አርጥርር
AAGTCTCTGG	AGTGGAGACT	CCTGGTCTAG	CATGACGACT	TCAGTTCACG
1060	1070	1090	1090	1100
1000	1010	1000	1090 * *	1100
* *	* *	* *	* *	* *
CATTGAGGTC	ATCATGTTGC	GCTCCAATGA	GTCCTTCACC	ATGGACGACA
			CAGGAAGTGG	
GTAACTCCAG	TAGTACAACG	CGAGGTTACT	CAGGAAGIGG	TACCTGCTGT
1110	1120	1130	1140	1150
4 4			4	
TGTCCTGGAC	CTGTGGCAAC	CAAGACTACA	AGTACCGCGT	CAGTGACGTG
A C A CC A C C T C	CACACCCTTC	CTTCTCATCT	TCATGGCGCA	CTCACTCCAC
ACAGGACCIG	GUCUCCGIIG	01101011101	101110000011	OTCHCIOCHC
1160	1170	1180	1190	1200
* *	* *	* *	1190 * *	* *
ACCAAAGCCG	GACACAGCCT	GGAGCTGATT	GAGCCCCTCA	TCAAGTTCCA
TGGTTTCGGC	CTGTGTCGGA	CCTCGACTAA	CTCGGGGAGT	AGTTCAAGGT
1210	1220	1230	1240	1250
* *	* *	* *	1240	* *
			GGAGGAGCAT	
CCACCCTGAC	TTCTTCGACT	TGAACGTACT	CCTCCTCGTA	CAGGACGAGT
1260	1270	1280	1290	1300
1200	1270	1200	1290 * *	100
TGGCCATCTG	CATCGTCTCC	CCAGATCGTC	CTGGGGTGCA	GGACGCCGCG
ACCGGTAGAC		CCTCTACCAC	CACCCCACCT	CCTCCCCCCC
	GTAGCAGAGG	GGTCTAGCAG	GACCCCACGT	CCTGCGGCGC
1310				
1310				
1310	1320	1330	1340	1350 * *
CTGATTGAGG	1320 * *	1330 * * CCGCCTGTCC	1340 * * AACACACTGC	1350 * * AGACGTACAT
CTGATTGAGG	1320 * *	1330 * * CCGCCTGTCC	1340 * * AACACACTGC	1350 * * AGACGTACAT
CTGATTGAGG	1320 * *	1330 * * CCGCCTGTCC	1340	1350 * * AGACGTACAT
CTGATTGAGG GACTAACTCC	1320 * * CCATCCAGGA GGTAGGTCCT	1330 * * CCGCCTGTCC GGCGGACAGG	1340 * * AACACACTGC TTGTGTGACG	1350 * * AGACGTACAT TCTGCATGTA
CTGATTGAGG GACTAACTCC 1360	1320 * * CCATCCAGGA GGTAGGTCCT 1370	1330  * * * * *CCGCCTGTCC GGCGGACAGG	1340 * * AACACACTGC TTGTGTGACG	1350  * AGACGTACAT TCTGCATGTA  1400
CTGATTGAGG GACTAACTCC	1320 * * CCATCCAGGA GGTAGGTCCT	1330  * * * * *CCGCCTGTCC GGCGGACAGG	1340 * * AACACACTGC TTGTGTGACG	1350 * * AGACGTACAT TCTGCATGTA
CTGATTGAGG GACTAACTCC 1360 * *	1320  * CCATCCAGGA GGTAGGTCCT  1370  *	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *	1340 * * AACACACTGC TTGTGTGACG 1390 * *	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA
CTGATTGAGG GACTAACTCC  1360 * * CCGCTGCCGC GGCGACGGCG	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC GTGGGCGGGG	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA CGGTTCTACT
CTGATTGAGG GACTAACTCC  1360 * * CCGCTGCCGC GGCGACGGCG	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC GTGGGCGGGG	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA CGGTTCTACT
CTGATTGAGG GACTAACTCC  1360 * * CCGCTGCCGC GGCGACGGCG  1410 * *	1320  * * CCATCCAGGA GGTAGGTCCT  1370  * * CACCCGCCCC GTGGGCGGGG  1420  * *	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT  1430  * *	1340  * * AACACACTGC TTGTGTGACG  1390  * * CCTGCTCTAT GGACGAGATA  1440  * *	1350  * *  AGACGTACAT TCTGCATGTA  1400  *  GCCAAGATGA CGGTTCTACT  1450  * *
CTGATTGAGG GACTAACTCC  1360 * * CCGCTGCCGC GGCGACGGCG  1410 * *	1320  * * CCATCCAGGA GGTAGGTCCT  1370  * * CACCCGCCCC GTGGGCGGGG  1420  * *	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT  1430  * *	1340  * * AACACACTGC TTGTGTGACG  1390  * * CCTGCTCTAT GGACGAGATA  1440  * *	1350  * *  AGACGTACAT TCTGCATGTA  1400  *  GCCAAGATGA CGGTTCTACT  1450  * *
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC GGCGACGGCG  1410  * TCCAGAAGCT	1320  * * CCATCCAGGA GGTAGGTCCT  1370  * * CACCCGCCC GTGGGCGGG  1420  * AGCCGACCTG	1330  * * CCGCCTGTCC GGCGGACAGG  1380  * * CGGGCAGCCA GCCCGTCGGT  1430  * CGCAGCCTCA	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA  1440  * *  ATGAGGAGCA	1350  * *  AGACGTACAT TCTGCATGTA  1400  *  GCCAAGATGA CGGTTCTACT  1450  *  CTCCAAGCAG
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC GGCGACGGCG  1410  * TCCAGAAGCT	1320  * * CCATCCAGGA GGTAGGTCCT  1370  * * CACCCGCCC GTGGGCGGG  1420  * AGCCGACCTG	1330  * * CCGCCTGTCC GGCGGACAGG  1380  * * CGGGCAGCCA GCCCGTCGGT  1430  * CGCAGCCTCA	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA  1440  * *  ATGAGGAGCA	1350  * *  AGACGTACAT TCTGCATGTA  1400  *  GCCAAGATGA CGGTTCTACT  1450  * *
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC GGCGACGGCG  1410  * TCCAGAAGCT	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC GTGGGCGGGG  1420  * AGCCGACCTG TCGGCTGGAC	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT  1430  * *  CGCAGCCTCA GCGTCGGAGT	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA  1440  *  ATGAGGAGCA TACTCCTCGT	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA CGGTTCTACT  1450  *  CTCCAAGCAG GAGGTTCGTC
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC GGCGACGGCG  1410  * TCCAGAAGCT AGGTCTTCGA	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC GTGGGCGGGG  1420  * AGCCGACCTG TCGGCTGGAC	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT  1430  * *  CGCAGCCTCA GCGTCGGAGT  1480	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA  1440  *  ATGAGGAGCA TACTCCTCGT	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA CGGTTCTACT  1450  *  CTCCAAGCAG GAGGTTCGTC
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC GGCGACGGCG  1410  * TCCAGAAGCT AGGTCTTCGA	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC GTGGGCGGGG  1420  * AGCCGACCTG TCGGCTGGAC	1330  * * CCGCCTGTCC GGCGGACAGG  1380  * * CGGGCAGCCA GCCCGTCGGT  1430  * * CGCAGCCTCA GCGTCGGAGT	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA  1440  *  ATGAGGAGCA TACTCCTCGT	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA CGGTTCTACT  1450  *  CTCCAAGCAG GAGGTTCGTC
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC GGCGACGGCG  1410  * TCCAGAAGCT AGGTCTTCGA  1460  *	1320  * * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC GTGGGCGGGG  1420  * AGCCGACCTG TCGGCTGGAC  1470  *	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT  1430  *  *  *  *  *  *  *  *  *  *  *  *  *	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA  1440  *  ATGAGGAGCA TACTCCTCGT  1490  * *	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA CGGTTCTACT  1450  *  CTCCAAGCAG GAGGTTCGTC  1500  * *
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC GGCGACGGCG  1410  * TCCAGAAGCT AGGTCTTCGA  1460  * TACCGCTGCC	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC GTGGGCGGGG  1420  * AGCCGACCTG TCGGCTGGAC  1470  * TCTCCTTCCA	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT  1430  *  CGCAGCCTCA GCGTCGGAGT  1480  *  GCCTGAGTGC	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA  1440  *  ATGAGGAGCA TACTCCTCGT  1490  *  AGCATGAAGC	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA CGGTTCTACT  1450  *  CTCCAAGCAG GAGGTTCGTC  1500  *  TAACGCCCCT
CTGATTGAGG GACTAACTCC  1360  * CCGCTGCCGC GGCGACGGCG  1410  * TCCAGAAGCT AGGTCTTCGA  1460  * TACCGCTGCC	1320  * CCATCCAGGA GGTAGGTCCT  1370  * CACCCGCCCC GTGGGCGGGG  1420  * AGCCGACCTG TCGGCTGGAC  1470  * TCTCCTTCCA	1330  * *  CCGCCTGTCC GGCGGACAGG  1380  * *  CGGGCAGCCA GCCCGTCGGT  1430  *  CGCAGCCTCA GCGTCGGAGT  1480  *  GCCTGAGTGC	1340  * *  AACACACTGC TTGTGTGACG  1390  * *  CCTGCTCTAT GGACGAGATA  1440  *  ATGAGGAGCA TACTCCTCGT  1490  *  AGCATGAAGC	1350  * *  AGACGTACAT TCTGCATGTA  1400  * *  GCCAAGATGA CGGTTCTACT  1450  *  CTCCAAGCAG GAGGTTCGTC  1500  * *

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1510 1520 1530

TGTGCTCGAA GTGTTTGGCA ATGAGATCTC CTGA ACACGAGCTT CACAAACCGT TACTCTAGAG GACT

FIGURE 8	TRANSO	CRIPT 11		
10	20	30	40	50
TGCGACCTTG	GCGGTGAGCC CGCCACTCGG	TGGGGACAGG		
60 *	70 *	80	90	100
	CCCGGCCCAA GGGCCGGGTT			
110	120	130	140	150 *
	GCGGTGTGTT CGCCACACAA			
160 *	170	180	190	200
	TGGTAGAAAG ACCATCTTTC			
210	220	230	240	250 *
CGTGCAGAAG GCACGTCTTC		TGAAGTGTCT ACTTCACAGA		
260	270 *	280	290 *	300
	TCCACTTACC AGGTGAATGG			
310	320	330	340	350
ACCGCCGGTC	CACTTCCCTG GTGAAGGGAC	CCTGACCCTG GGACTGGGAC	GAGACTTTGA CTCTGAAACT	GGCCTTGCAC
		-		o ArgAsnVal>
360	370	380	, 390 *	*
CCCCGGATCT GGGGCCTAGA ProArgIle	CACCCCACAC	TGGAGACCGA ACCTCTGGCT s GlyAspArg	CGGTGACCGA	TTCACTTCAA AAGTGAAGTT PheHisPheAsn>
410			440	450
* TGCTATGACC	* TGTGAAGGCT ACACTTCCGA	* GCAAAGGCTT	* CTTCAGGCGA	AGCATGAAGC
AlaMetThr	CysGluGly	CysLysGlyPh	e PheArgArg	SerMetLys>
460	*	*	490	500 *
CCTTCCGTGA	ATTCACCTGC TAAGTGGACG u PheThrCys	GGGAAGTTGC	CCCTGACGGC	CATCACCAAG GTAGTGGTTC g IleThrLys>

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510 520 530 540 550 * GACAACCGAC GCCACTGCCA GGCCTGCCGG CTCAAACGCT GTGTGGACAT CTGTTGGCTG CGGTGACGGT CCGGACGGCC GAGTTTGCGA CACACCTGTA AspAsnArg ArgHisCysGln AlaCysArg LeuLysArg CysValAspIle> 570 580 590 600 CGGCATGATG AAGGAGTTCA TTCTGACAGA TGAGGAAGTG CAGAGGAAGC GCCGTACTAC TTCCTCAAGT AAGACTGTCT ACTCCTTCAC GTCTCCTTCG GlyMetMet LysGluPhe IleLeuThrAsp GluGluVal GlnArgLys> 620 610 630 640 650 GGGAGATGAT CCTGAAGCGG AAGGAGGAGG AGGCCTTGAA GGACAGTCTG CCCTCTACTA GGACTTCGCC TTCCTCCTCC TCCGGAACTT CCTGTCAGAC ArgGluMetIle LeuLysArg LysGluGlu GluAlaLeuLys AspSerLeu> 680 660 690 700 CGGCCCAAGC TGTCTGAGGA GCAGCAGCGC ATCATTGCCA TACTGCTGGA GCCGGGTTCG ACAGACTCCT CGTCGTCGCG TAGTAACGGT ATGACGACCT ArgProLys LeuSerGluGlu GlnGlnArg IleIleAla IleLeuLeuAsp> 740 750 710 720 730 CGCCCACCAT AAGACCTACG ACCCCACCTA CTCCGACTTC TGCCAGTTCC GCGGGTGGTA TTCTGGATGC TGGGGTGGAT GAGGCTGAAG ACGGTCAAGG AlaHisHis LysThrTyr AspProThrTyr SerAspPhe CysGlnPhe> 790 800 760 770 780 GGCCTCCAGT TCGTGTGAAT GATGGTGGAG GGAGCCATCC TTCCAGGCCC CCGGAGGTCA AGCACACTTA CTACCACCTC CCTCGGTAGG AAGGTCCGGG ArgProProVal ArgValAsn AspGlyGly GlySerHisPro SerArgPro> 810 820 830 840 850 AACTCCAGAC ACACTCCCAG CTTCTCTGGG GACTCCTCCT CCTCCTGCTC TTGAGGTCTG TGTGAGGGTC GAAGAGACCC CTGAGGAGGA GGAGGACGAG AsnSerArg HisThrProSer PheSerGly AspSerSer SerSerCysSer> 860 870 890 AGATCACTGT ATCACCTCTT CAGACATGAT GGACTCGTCC AGCTTCTCCA TCTAGTGACA TAGTGGAGAA GTCTGTACTA CCTGAGCAGG TCGAAGAGGT AspHisCys IleThrSer SerAspMetMet AspSerSer SerPheSer> 940 950 910 920 930 ATCTGGATCT GAGTGAAGAA GATTCAGATG ACCCTTCTGT GACCCTAGAG TAGACCTAGA CTCACTTCTT CTAAGTCTAC TGGGAAGACA CTGGGATCTC AsnLeuAspLeu SerGluGlu AspSerAsp AspProSerVal ThrLeuGlu> 1000 990 980 960 CTGTCCCAGC TCTCCATGCT GCCCCACCTG GCTGACCTGG TCAGTTACAG GACAGGGTCG AGAGGTACGA CGGGGTGGAC CGACTGGACC AGTCAATGTC LeuSerGln LeuSerMetLeu ProHisLeu AlaAspLeu ValSerTyrSer>

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*	1020	1030	1040	1050 *
CATCCAAAAG GTAGGTTTTC IleGlnLys	GTCATTGGCT CAGTAACCGA VallleGly I	AACGATTCTA	CTATGGTCCT	AAGTCTCTGG
1060	1070	1080	1090 *	1100
AGTGGAGACT	GGACCAGATC CCTGGTCTAG AspGlnIle	CATGACGACT	TCAGTTCACG	CATTGAGGTC GTAACTCCAG a IleGluVal>
1110	1120	1130	1140	1150 *
TAGTACAACG	GCTCCAATGA CGAGGTTACT \rgSerAsnGlu	CAGGAAGTGG	TACCTGCTGT	
1160 *	1170	1180	1190	1200
GACACCGTTG	CAAGACTACA GTTCTGATGT GlnAspTyr I	TCATGGCGCA	GTCACTGCAC	TGGTTTCGGC
1210	1220	1230	1240	1250
CTGTGTCGGA	GGAGCTGATT CCTCGACTAA	CTCGGGGAGT	AGTTCAAGGT	CCACCCTGAC
GlyHisSerLe	ı GluLeuIle	GluProLeu	ITeLyspheGI	n valGlyLeu>
GlyHisSerLet	1270	1280	IIeLysPheGI 1290 *	1300
GlyHisSerLet  1260  *  AAGAAGCTGA TTCTTCGACT	1270 * ACTTGCATGA TGAACGTACT	1280 * GGAGGAGCAT CCTCCTCGTA	1290 * GTCCTGCTCA CAGGACGAGT	1300 * TGGCCATCTG
GlyHisSerLet  1260  *  AAGAAGCTGA TTCTTCGACT	1270 * ACTTGCATGA TGAACGTACT	1280 * GGAGGAGCAT CCTCCTCGTA	1290 * GTCCTGCTCA CAGGACGAGT	1300 * TGGCCATCTG ACCGGTAGAC
GlyHisSerLet  1260  *  AAGAAGCTGA TTCTTCGACT LysLysLeu  1310  *  CATCGTCTCC GTAGCAGAGG	1270  * ACTTGCATGA TGAACGTACT AsnLeuHisGlu 1320 * CCAGATCGTC GGTCTAGCAG	1280  * GGAGGAGCAT CCTCCTCGTA GluGluHis 1330 * CTGGGGTGCA GACCCCACGT	1290  * GTCCTGCTCA CAGGACGAGT ValLeuLeu  1340  * GGACGCCGCG CCTGCGGCGCC	1300 * TGGCCATCTG ACCGGTAGAC MetAlalleCys> 1350 * CTGATTGAGG GACTAACTCC
GlyHisSerLet  1260  *  AAGAAGCTGA TTCTTCGACT LysLysLeu  1310  *  CATCGTCTCC GTAGCAGAGG IleValSer	1270  * ACTTGCATGA TGAACGTACT AsnLeuHisGlu 1320  * CCAGATCGTC GGTCTAGCAG ProAspArg	1280  * GGAGGAGCAT CCTCCTCGTA GluGluHis 1330  * CTGGGGTGCA GACCCCACGT ProGlyValGl	1290  * GTCCTGCTCA CAGGACGAGT ValLeuLeu 1 1340  * GGACGCCGCG CCTGCGGCGC n AspAlaAla	1300  * TGGCCATCTG ACCGGTAGAC MetAlalleCys> 1350  * CTGATTGAGG GACTAACTCC LeulleGlu>
GlyHisSerLet  1260  * AAGAAGCTGA TTCTTCGACT LysLysLeu  1310  * CATCGTCTCC GTAGCAGAGG IleValser  1360  *	1270  * ACTTGCATGA TGAACGTACT AsnLeuHisGlu 1320  * CCAGATCGTC GGTCTAGCAG ProAspArg 1 1370  *	1280  * GGAGGAGCAT CCTCCTCGTA GluGluHis 1330  * CTGGGGTGCA GACCCCACGT ProGlyValGli	1290  * GTCCTGCTCA CAGGACGAGT ValLeuLeu l  1340  * GGACGCCGCG CCTGCGGCGC n AspAlaAla  1390 *	1300  * TGGCCATCTG ACCGGTAGAC MetAlalleCys>  1350  * CTGATTGAGG GACTAACTCC LeulleGlu>  1400  *
GlyHisSerLet  1260  * AAGAAGCTGA TTCTTCGACT LysLysLeu  1310  * CATCGTCTCC GTAGCAGAGG IleValser  1360  * CCATCCAGGA GGTAGGTCCT	1270  * ACTTGCATGA TGAACGTACT AsnLeuHisGlu 1320  * CCAGATCGTC GGTCTAGCAG ProAspArg 1370  * CCGCCTGTCC GGCGGACAGG	1280  * GGAGGAGCAT CCTCCTCGTA GluGluHis 1330  * CTGGGGTGCA GACCCCACGT ProGlyValGli 1380  * AACACACTGC	1290  * GTCCTGCTCA CAGGACGAGT ValLeuLeu  1340  * GGACGCCGCG CCTGCGGCGC n AspAlaAla  1390  * AGACGTACAT TCTGCATGTA	1300  * TGGCCATCTG ACCGGTAGAC MetAlalleCys>  1350  * CTGATTGAGG GACTAACTCC LeulleGlu>  1400  *
GlyHisSerLet  1260  * AAGAAGCTGA TTCTTCGACT LysLysLeu  1310  * CATCGTCTCC GTAGCAGAGG IleValser  1360  * CCATCCAGGA GGTAGGTCCT	1270  * ACTTGCATGA TGAACGTACT AsnLeuHisGlu 1320  * CCAGATCGTC GGTCTAGCAG ProAspArg 1370  * CCGCCTGTCC GGCGGACAGG	1280  * GGAGGAGCAT CCTCCTCGTA GluGluHis 1330  * CTGGGGTGCA GACCCCACGT ProGlyValGli 1380  * AACACACTGC	1290  * GTCCTGCTCA CAGGACGAGT ValLeuLeu  1340  * GGACGCCGCG CCTGCGGCGC n AspAlaAla  1390  * AGACGTACAT TCTGCATGTA	1300  * TGGCCATCTG ACCGGTAGAC MetAlalleCys>  1350  * CTGATTGAGG GACTAACTCC LeulleGlu>  1400  * CCGCTGCCGC GGCGACGCG
GlyHisSerLet  1260  * AAGAAGCTGA TTCTTCGACT LysLysLeu // 1310  * CATCGTCTCC GTAGCAGAGG IleValSer  1360  * CCATCCAGGA GGTAGGTCCT AlaIleGlnAs; 1410  * CACCCGCCCC GTGGGCGGGG	1270  * ACTTGCATGA TGAACGTACT AsnLeuHisGlu 1320 * CCAGATCGTC GGTCTAGCAG ProAspArg  1370 * CCGCCTGTCC GGCGGACAGG PArgLeuSer 1420 * CGGGCAGCCA CGGCCGGCCAGCCA GCCCGTCGGTCC	GGAGGAGCAT CCTCCTCGTA GIUGIUHIS 1330  CTGGGGTGCA GACCCCACGT ProGlyValGI  AACACACTGC TTGTGTGACG ASnThrLeu  1430  CCTGCTCTAT GGACGAGATA	1290  * GTCCTGCTCA CAGGACGAGT ValLeuLeu  1340  GGACGCCGCG CCTGCGGCGC n AspAlaAla  1390  * AGACGTACAT TCTGCATGTA GINThrTyrIl  1440  GCCAAGATGA CGGTTCTACT	1300  * TGGCCATCTG ACCGGTAGAC MetAlalleCys>  1350  * CTGATTGAGG GACTAACTCC LeulleGlu>  1400  * CCGCTGCCGC GGCGACGCGC e ArgCysArg>
GlyHisSerLet  1260  * AAGAAGCTGA TTCTTCGACT LysLysLeu // 1310  * CATCGTCTCC GTAGCAGAGG IleValSer  1360  * CCATCCAGGA GGTAGGTCCT AlaIleGlnAs; 1410  * CACCCGCCCC GTGGGCGGGG	1270  * ACTTGCATGA TGAACGTACT AsnLeuHisGlu 1320 * CCAGATCGTC GGTCTAGCAG ProAspArg  1370  * CCGCCTGTCC GGCGGACAGG PArgLeuSer 1420 * CGGGCAGCCA GCCCGTCGGT ProGlySerHi	GGAGGAGCAT CCTCCTCGTA GIUGIUHIS 1330  CTGGGGTGCA GACCCCACGT PROGLYVALGI  AACACACTGC TTGTGTGACG ASNTHRLEU  1430  * CCTGCTCTAT GGACGAGATA S LeuLeuTyr	1290  * GTCCTGCTCA CAGGACGAGT ValLeuLeu  1340  GGACGCCGCG CCTGCGGCGC  ASPAlaAla  1390  * AGACGTACAT TCTGCATGTA GInThrTyrIl  1440  * GCCAAGATGA CGGTTCTACT AlaLysMet	TGGCCATCTG ACCGGTAGAC MetAlalleCys>  1350  CTGATTGAGG GACTAACTCC LeuIleGlu>  1400  * CCGCTGCCGC GGCGACGGCG e ArgCysArg>  1450  * TCCAGAAGCT AGGTCTTCGA IleGlnLysLeu>

1510 1520 1530 1540 1550

TCTCCTTCCA GCCTGAGTGC AGCATGAAGC TAACGCCCCT TGTGCTCGAA AGAGGAAGGT CGGACTCACG TCGTACTTCG ATTGCGGGGA ACACGAGCTT LeuSerPheGln ProGluCys SerMetLys LeuThrProLeu ValLeuGlu>

1560 1570

GTGTTTGGCA ATGAGATCTC CTGA CACAAACCGT TACTCTAGAG GACT ValPheGly AsnGlulleSer ***>

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## DECLARATION, POWER OF ATTORNEY AND PETITION

As a balow named inventor, I hereby declars that:

My residence, past office and civizenship are as stated below next to tay name.

I believe I am the original, first and sole inventor (if only one mane is listed below) or an original first and joint inventor (if plure) nomes ore listed below] of the subject matter claimed and for winds a patent is sought on the invention entitled:

#### ISOFORMS OF THE HUMAN VITAMIN D RECEPTOR

the specification of Which

🗋 is attached horsto. 🕱 was filed on 29 September 1998 as Application No. PCT/AUSE/00817 and was amended on (if appliesble).

I hereby state that I have reviewed and understand the consents of the above identified specification, including the claims, as nineddad by any amendment referred to above.

I neknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, Section 1.56(a)

History claim foreign priority ben-fits under Title 35, United States Code, Section 119 of any foreign application(s) for parant or inventor's certificate listed below and have also identified below any fereign application for patent or inventor's scruticate having a filling date before that of the application on which priority is claimed:

	Princ Fureign Appli	ention(z)	. Prioris	Claimed
PO 9500	Austialia	29 September 1997	<u> </u>	
Number	[Country]	[Pay/Month/Yout Filed]	Yes	No

I hereby claim the benefit under Tirle 35, United States Code, Section 120 of any United State; application(s) listed below and, insufar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the commer provided by the first paragraph of Title 35; United States Code, Section 112, I seknow edge the dury to disclose material information as defined in Title 37, Code of Federal Regulations, Section (1.55(x) which occurred between the filling date of the prior application and the national or PCT international fifing date of this application.

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(Annication Serial (to)	[Filing Date]	Sentus: parantol, pending, abandonedi

I hereby decime that all statement a made herein of my own knowledge are true and that all at rements made on information and belief are believed to be true; and further that these stotements were used with the knowledge; that within take statements and tha like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such will'd false statements may jeopardize the validity of the application or any patent issued thereon.

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with full power of substitution and revocation, to prosecute this opplication and to transact all business in the Patent and Trademark Office connected therewith, and all future correspondence should be addressed to trem.

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## DECLARATION, FOWER OF ATTORNEY AND PETITION

As a below numbed in enter, I hereby declare that

My registance, post office and citizenship are as stated below acut to my name,

I believe I am the original. Het and tole inventor (if only one name is listed below) or an original in at and joint inventor (if plurat nonjes and listed below) of the subject matter claimed and for which a patent is sought on the investigate entitled:

#### ISOFORMS OF THE HUMAN VITAMIN D RECEPTIOR

the specification of Wi	delt.	! •
is attached hereto	was filed on 29 September 1998 as Application No. PCT/Au98/00817	भारत <i>प्रकृत भारता देवचे वर</i>
(if applicable).	-	:

I hereby state that I have reviewed and understand the consents of the above identified specification. Including the claims, as unended by any unendment referred to above.

Lacknowledge the duty to disclose hubbroatton which is material to the examination of this applies from in accordance with Title 37. Cade of Federal Regulations, Section 1.56(a)

History claim foreign priority benefits under Title 35. United States Code, Section 119 of any for ign application(s) for parent or inventor's carlificate listed below and have also identified below any foreign application for pateot or inventor's carlificate having a filling date before that of the application on which priority is claimed:

	Priority	C almed		
PO 9500	Australia	29 September 1997	<b>3</b>	
[Number]	Country	Day/Month/Year Filed	Yes	No

! hereby claim the benefit under Title 35. United States Code, Section 120 of any United States application(z) listed believe and involve the subject matter of each of the eletins of this application is not disclosed in the prior littled States application is not disclosed in the prior littled States application is not disclosed in the prior littled States application is not disclosed in the prior littles are not disclosed in the inclusion of the little states of the day to disclose metable nationalists and fine in Title 37. Code of Federal Regulations, Section 1.56(a) which accurred hereven the fillingidate of the micro application and the national or PLT international filing date of this application.

Marie Marie Anna Marie M		
(Application Serial pd)	Filing Dotal	[Status: patented, sending, changaned]

I hereby declare that all statement a mule herein of my own knowledge are true and that all statements made on information and build are believed to be true; and further that these stotements were ande with the knowledge that withit false statements and the like so made are publishable by they of imprisonment or both, under Section 1001 of Title 18 of the United States Code and that such will'd false sistements may jeopardize the validity of the application or any patent issued thereon.

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with full power of substitution and revocation, to prosecute this application and to transact all business in the Fatcht and Fractionarie Office connected therewith and all future correspondence should be addressed to them.

full name of sule as first inventor: t	Jroffs, lánda Adar
Inventor's Signature	DateDate
Chizenship: Australian	neville. New South Wales 2043, Australia 201. Erskinovilla. New South Wales 2048, Austrolia

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# DECLARATION, POWER OF ATTORNEY AND PETITION

Ac a below unimed invaluer, I hereby declare that:

My tosidence, Just office and chizenship are as stated below next to my heart,

I believe I am the original. first and sale inventor (if only one name is listed below) or an original tirst and joint inventor (if plural names are listed below) of the subject mater claimed and for which a paints is sought on the investion cutified;

### ISOFORMS OF THE HUMAN VITAMIN D RECEPT OR

the specification of wi	nich	ب دلد
is attached hereto	nich 💢 was filed an IS September 1998 as Application No. PCT/AU9E/00817 and was airt	puoda da
(if applicable).	·	:

I hereby state that I have reviewed and understand the concents of the above identified excellination, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to dischase information which is material to the examination of this applies tion in accordance with Title 37. Code of Federal Regulations, Section 1,56(2)

Thereby slaim foreign priority benefits under l'ille 35, United States Code, Section 119 of any fort ign application(s) for patent or inventor's certificate listed below and have also identified below any lessing application for patent or inventor's certificate baying a filling date before that of the application on which priority is claimed:

	Priority	C) elmad		
TO 9500	Australia	29 September 1997		Ī.Д
[Number]	[Caun'sy]	[Day/Month/Your Filed]	Yez	No

I brokely claim the benefit moder Title 14. Thitsel States Code, Section 120 of any United States as plication(a) histed below and, insular as the subject matter of each of the chains of this application is not disclosed in the prior I filed States application in the matter provided by the first paragraph of Title 35. United States Code, Section 112, I acknowledge the duty to displace insterial information as defined in Title 37, Code of Pederal Regulations, Section 1.56(a) which occurred between the filing-late of the prior application and the national or PC f international filling date of this application.

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(Application Serial hul	[Filling Date]	(Stutus: patented, page	ing, aliantinneil]

I hereby deplate that all statement a mode hereix of the over knowledge are true and that all statements made on information and bulled are believed to be true; and further that dues and encourage were made with the knowledge that within false statements and the like so made are punishable by time or imprisonment, or both, under Section 1601 of Title 18 of the United States Code and that such willful false statements may isopardize the validity of the application or any patent instead the mon.

> ROTHWELL FIGG ERNST & KURZ Columbia Square Suite 701, Bust Towns Washington, District of Columbia, 20104 United States of America

with full power of substitution and envecation, to proceed this application and to transact all but most in t	he Patent and
Produces to Office connected therewith, and all future unrespondence should be addressed to them.	<b>:</b>

Full name of sole or first invastor: CROFFS, Linda Anna	
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JUNIZANT

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# DECLARATION, POWER OF ATTORNEY AND PETITION

As a below named inventor, I hereby declare that:

My residence, post office and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original first and joint inventor (if plural names are listed below) of the subject matter claimed and for which a patent is sought on the invention entitled:

### ISOFORMS OF THE HUMAN VITAMIN D RECEPTOR

the specification of wh	nich	÷
is attached hereto	was filed on 29 September 1998 as Application No. PCT/AU98/00(17 and was amen	aded or
((Capplicable)	*	•

I hereby siste that I have reviewed and understand the consents of the above identified specification, including the claims, as unended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37. Code of Federal Regulations, Section 1.56(a)

I hereby claim foreign priority benefits under Title 35, United States Code, Section 119 of any loreign application(s) for patent or inventor's cartificate histories application for patent of the application on which priority is claimed:

Princ Foreign Application(s)			Priority Claimed	
PO 9500	Australia	29 September 1997	区	
[Number]	[Country]	[Day/Month/Yest Filed]	Yes	No

I hereby claim the benefit under Title 35, United States Code, Section 120 of any United State; application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35. United States Code, Section 112, Lecknow edge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, Section 1.56(a) which occurs d between the tiling date of the prior application and the national or PCT international filing date of this application.

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[Application Serial no]	(Filing Date)	(Status: patente l, pending, abandoned)

I hereby declare that all statement's made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that wifful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such wilful false statements may jeopardize the validity of the application or any patent issued thereon.

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with full power of substitution and revocation, to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith, and all future correspondence should be addressed to hear.

Full name of solo or first inventor: CROFTS, Linda Anne					
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